Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	68389	diabet\$	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L2	33009	phenylalan\$	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L3	5049	diabet\$ and phenylalan\$	USPAT; EPO; JPO; DERWENT	OR .	OFF	2005/04/27 06:01
L4	1450	PPAR	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L5	445	(562/431).CCLS.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L6	383	(562/445).CCLS.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L7	0	("l2andl11").PN.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L8	573	(514/538).CCLS.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L9	0	("dibutylbenzene").PN.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L10	0	"5059736".URPN.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L11	5962	integrin\$	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L12	5049	diabet\$ and (diabet\$ and phenylalan\$)	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L13	2172	diabet\$ and integrin\$	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L14	1649	NIDDM	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01

L15	1059	(562/426).CCLS.	USPAT; USOCR; EPO; JPO;	OR	OFF	2005/04/27 06:01
L16	910	(514/563).CCLS.	DERWENT USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L17	58	(diabet\$ and phenylalan\$) and PPAR	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L18	27	diabet\$ and (("562/445").CCLS.)	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L19	3	"9935163".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L20	3	diabet\$ and (("562/446").CCLS.)	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L21	3	"9943642".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L22	5	"9906431".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L23	2	"9622966".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L24	3	"9515973".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L25	25	diabet\$ and (("562/431").CCLS.)	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L26	2	("5321181").PN.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L27	2	("4849569").PN.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L28	86	diabet\$ and (("514/538").CCLS.)	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L29	2	("5055627").PN.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01

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L30	2	("4950834").PN.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L31	6	"748784".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L32	2	("5059736").PN.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L33	2	di-n-butylbenzene	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L34	7	"9935163"	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L35	2	"5158959".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L36	12	PPAR and (diabet\$ and integrin\$)	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L37	17	NIDDM and integrin\$	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L38	2	"11140079"	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 06:01
L39	166	dibutylbenzene	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L40	204	(562/446).CCLS.	USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L41	4	("5216167").PN.	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L42	5	"655562".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L43	2	"6555562".pn.	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L44	1	"4987132".PN.	USPAT; USOCR	OR	ON	2005/04/27 06:01

L45	1	"5164372".PN.	USPAT; USOCR	OR	ON	2005/04/27 06:01
L46	1	"5260277".PN.	USPAT; USOCR	OR	ON	2005/04/27 06:01
L47	1	"5296486".PN.	USPAT; USOCR	OR	ON	2005/04/27 06:01
L48	1	"5399585".PN.	USPAT; USOCR	OR	ON	2005/04/27 06:01
L49	1	"6093696".PN.	USPAT; USOCR	OR	ON	2005/04/27 06:01
L50	654	L5 or L6	USPAT; EPO; JPO; DERWENT	OR	OFF	2005/04/27 06:01
L51	26	L50 and L3	USPAT; EPO; JPO; DERWENT	OR .	OFF	2005/04/27 06:01
L52	11	"9736859"	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 06:24
L53	10	"0710657"	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 06:48
L54	169130	pharmaceutically adj acceptable	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 06:48
L55	197582	hexane	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 06:48
L56	627	I54 same I55	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 07:30
L57	0	("2005/0075377").URPN.	USPAT	OR	ON	2005/04/27 06:49
L58	0	"l6713514".pn.	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 07:30
L59	2	"6713514".pn.	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	ON	2005/04/27 07:30

	Туре	L #	Hits	Search Text	DBs	Time Stamp
1	BRS	L1	68389	diabet\$	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
2	BRS	L2	33009	phenylalan\$	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
3	BRS	L3	5049	diabet\$ and phenylalan\$	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
4	BRS	L4	1450	PPAR		2005/04/27 06:01
5	IS&R	L5	445	(562/431).CCLS.		2005/04/27 06:01
6	IS&R	L6	383	(562/445).CCLS.		2005/04/27 06:01

	Comments	Error Definition	Err ors
1			
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	Туре	L#	Hits	Search Text	DBs	Time Stamp
7	IS&R	L7	0	("l2andl11").PN.	USPAT; USOCR; EPO; JPO; DERWEN T	2005/04/27 06:01
8	IS&R	L8	573	(514/538).CCLS.	USPAT; USOCR; EPO; JPO; DERWEN	2005/04/27 06:01
9	IS&R	L9	0	("dibutylbenzene").PN.	USPAT; USOCR; EPO; JPO; DERWEN	2005/04/27 06:01
10	BRS	L10	0	"5059736".URPN.	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
11	BRS	L11	5962	integrin\$	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
12	BRS	L12	15 1 1 Z1 G	diabet\$ and (diabet\$ and	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01

	Comments	Error Definition	Err
7			
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9			
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12			

	Туре	L #	Hits	Search Text	DBs	Time Stamp
13	BRS	L13	2172	diabet\$ and integrin\$	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
14	BRS	L14	1649	NIDDM	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
15	IS&R	L15	1059	(562/426).CCLS.		2005/04/27 06:01
16	IS&R	L16	910	(514/563).CCLS.		2005/04/27 06:01
17	BRS	L50	654		USPAT; EPO; JPO; DERWEN	2005/04/27 06:01
18	BRS	L44	1	"4987132".PN.		2005/04/27 06:01
19	BRS	L45	1	"5164372".PN.		2005/04/27 06:01
20	BRS	L46	1	"5260277".PN.		2005/04/27 06:01

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	Туре	L #	Hits	Search Text	DBs	Time Stamp
21	BRS	L47	1	"5296486".PN.	USPAT; USOCR	2005/04/27 06:01
22	BRS	L48	1	"5399585".PN.	USPAT; USOCR	2005/04/27 06:01
23	BRS	L49	1	"6093696".PN.	USPAT; USOCR	2005/04/27 06:01
24	BRS	L17	58	(diabet\$ and phenylalan\$) and PPAR	USPAT; EPO; JPO; DERWEN	2005/04/27 06:01
25	BRS	L18	27	diabet\$ and (("562/445").CCLS.)	USPAT; EPO; JPO; DERWEN	2005/04/27 06:01
26	BRS	L19	3	"9935163".pn.	USPAT; EPO; JPO; DERWEN	2005/04/27 06:01
27	BRS	L20	3	diabet\$ and (("562/446").CCLS.)	USPAT; EPO; JPO; DERWEN	2005/04/27 06:01
28	BRS	L21	3	"9943642".pn.	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
29	BRS	L22	5	"9906431".pn.	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01

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	Туре	L #	Hits	Search Text	DBs	Time Stamp
30	BRS	L23	2	"9622966".pn.	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
31	BRS	L24	3	"9515973".pn.	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
32	BRS	L25	25	diabet\$ and (("562/431").CCLS.)		2005/04/27 06:01
33	IS&R	L26	2	("5321181").PN.		2005/04/27 06:01
34	IS&R	L27	2	("4849569").PN.		2005/04/27 06:01
35	BRS	L28	86	diabets and (("514/538").CCLS.)	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01

	Comments	Error Definition	Err
30			
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35			

	Туре	L#	Hits	Search Text	DBs	Time Stamp
36	IS&R	L29	2	("5055627").PN.		2005/04/27 06:01
37	IS&R	L30	2	("4950834").PN.		2005/04/27 06:01
38	BRS	L31	6	"748784".pn.		2005/04/27 06:01
39	IS&R	L32	2	("5059736").PN.		2005/04/27 06:01
40	BRS	L33	2	di-n-butylbenzene	. I D{ } •	2005/04/27 06:01
41	BRS	L34	7	"9935163"	k I D() •	2005/04/27 06:01

	Comments	Error Definition	Err
36			
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41	,		

	Туре	L #	Hits	Search Text	DBs	Time Stamp
42	BRS	L35	2	"5158959".pn.	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
43	BRS	L36	11 /	PPAR and (diabet\$ and integrin\$)	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
44	BRS	L37	17	NIDDM and integrin\$	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
45	BRS	L38	2	"11140079"	US- PGPUB; USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
46	IS&R	L41	4	("5216167").PN.		2005/04/27 06:01
47	BRS	L42	5	"655562".pn.		2005/04/27 06:01

	Comments	Error Definition	Err ors
42			
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	Туре	L #	Hits	Search Text	DBs	Time Stamp
48	BRS	L43	2	"6555562".pn.	USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
49	BRS	L51	26			2005/04/27 06:01
50	BRS	L39	166		USPAT; EPO; JPO; DERWEN T	2005/04/27 06:01
51	IS&R	L40	204	(562/446).CCLS.		2005/04/27 06:01
52	BRS	L52	11	"9736859"		2005/04/27 06:24
53	BRS	L53	10	"0710657"	IH: P(I *	2005/04/27 06:48

	Comments	Error Definition	Err ors
48			
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	Туре	L #	Hits	Search Text	DBs	Time Stamp
54	BRS	L54		pharmaceutically adj acceptable		2005/04/27 06:48
55	BRS	L55	197582	hexane		2005/04/27 06:48
56	BRS	L57	0	("2005/0075377").URPN.	USPAT	2005/04/27 06:49
57	BRS	L56	627	154 same 155	US- PGPUB; USPAT; EPO; JPO; DERWEN	2005/04/27 07:30
58	BRS	L58	0		US- PGPUB; USPAT; EPO; JPO; DERWEN	2005/04/27 07:30
59	BRS	L59	2	"6713514".pn.	US- PGPUB; USPAT; EPO; JPO; DERWEN	2005/04/27 07:30

	Comments	Error Definition	Err ors
54		· *	
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               AND CURRENT DISCOVER FILE IS DATED 11 AUGUST 2004
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0.21

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SINCE FILE TOTAL ENTRY SESSION 0.42 0.63

FULL ESTIMATED COST

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SINCE FILE TOTAL ENTRY SESSION 0.42 0.63

FULL ESTIMATED COST

L1 STRUCTURE UPLOADED

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L1 HAS NO ANSWERS

L1

STR

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=> search l1 exact full FULL SEARCH INITIATED 06:24:53 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 100 TO ITERATE

100.0% PROCESSED 100 ITERATIONS

4 ANSWERS

SEARCH TIME: 00.00.01

L2 4 SEA EXA FUL L1

=> d scan

L2 4 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN IN 1,4-Benzenedi(acetic-d2) acid, diethyl ester (9CI) MF C14 H14 D4 O4

$$\begin{array}{c|c} & \circ & \circ \\ \vdots & \vdots & \vdots \\ c D_2 - C - OEt \end{array}$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):4

L2 4 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 1,4-Benzene-2,3,5,6-d4-diacetic acid, diethyl ester (9CI)

MF C14 H14 D4 O4

$$\begin{array}{c|c}
D & O \\
CH_2-C-OEt
\end{array}$$
Eto-C-CH₂

L2 4 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN Poly[oxy-1,4-butanediyloxy(1-oxo-1,2-ethanediyl)-1,4-phenylene(2-oxo-1,2-ethanediyl)] (9CI)

MF (C14 H16 O4)n

CI PMS

RELATED POLYMERS AVAILABLE WITH POLYLINK

L2 4 ANSWERS REGISTRY COPYRIGHT 2004 ACS on STN

IN 1,4-Benzenediacetic acid, diethyl ester (9CI)

MF C14 H18 O4

CI COM

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

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FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION 53.09 53.30

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 12

L3 21 L2

=> 12/prep

21 L2

3212539 PREP/RL

L4

11 L2/PREP

(L2 (L) PREP/RL)

=> d 14 5-11 ti fbib abs

L4 ANSWER 5 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN

TI Palladium complex-catalyzed carboalkoxylation of bis(chloromethyl)arenes

AN 1988:454438 CAPLUS

DN 109:54438

TI Palladium complex-catalyzed carboalkoxylation of bis(chloromethyl)arenes

AU Kobayashi, Toshiaki; Abe, Fujio; Tanaka, Masato

CS Natl. Chem. Lab. Ind., Yatabe, 305, Japan

SO Journal of Molecular Catalysis (1988), 45(1), 91-109

CODEN: JMCADS; ISSN: 0304-5102

DT Journal

LA English

OS CASREACT 109:54438

GΙ

$$RCH_2$$
 CH_2R CH_2

Carboalkoxylation of 4-ClCH2C6H4CH2Cl with ROH (R = Me, Et, Me2CH, Me3C, AB Ph) and CO in the presence of PdCl2(PPh3)2 and N, N-dicyclohexylmethylamine gave diesters 4-RO2CCH2C6H4CH2CO2R as the major products. A similar reaction of 8 other bis(chloromethyl)arenes, e.g. I, II, and III (R = Cl), with MeOH and CO gave the corresponding diesters I, II, and III (R =CO2Me). Reaction parameters, such as auxiliary base, palladium complex catalyst, and solvent, were found to significantly affect the selectivity for diester formation.

III

II

- L4ANSWER 6 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- ΤI Syntheses of arenediacetic esters and acetonyl-substituted arylacetic esters by means of Friedel-Crafts reaction with α -acyl- α chlorosulfides
- 1986:590606 CAPLUS AN
- DN 105:190606
- ΤI Syntheses of arenediacetic esters and acetonyl-substituted arylacetic esters by means of Friedel-Crafts reaction with α -acyl- α chlorosulfides
- ΑU Ishibashi, Hiroyuki; Ikeda, Masazumi; Choi, Hong Dae; Nakagawa, Hiroko; Ueda, Yuko; Tamura, Yasumitsu
- Kyoto Pharm. Univ., Kyoto, 607, Japan CS
- SO Chemical & Pharmaceutical Bulletin (1985), 33(12), 5310-15 CODEN: CPBTAL; ISSN: 0009-2363
- DTJournal
- English LΑ
- os CASREACT 105:190606

GI

CHMeCO2Me

- AB Friedel-Crafts reaction of 2,5-R2C6H3CO2Et (R = H, Me) with $\alpha\text{-chloro-}\alpha\text{-}\text{(methylthio)}$ acetate (I) in the presence of SnCl4 gave the $\alpha\text{-methylthio-1}$,4-benzenediacetates II. The reactions of biphenyl, Ph2CH2, and Ph2O with excess I gave III (Rl = MeS, X = bond, CH2, O, resp.). Desulfurization of II (R = H, Me) and these III gave 2,5-R2C6H2(CH2CO2Et)2-1,4 and III (Rl = H). Me 4-(2-oxopropyl)phenylacetate was prepared by reaction of Me phenylacetate with $\alpha\text{-chloro-}\alpha\text{-}\text{-}\text{(methylthio)}$ acetone (IV) followed by desulfurization of the resulting product. Me 2-(2-furyl)propionate treated with IV in the presence of ZnCl2 gave furan V (Rl = MeS), desulfurization of which gave V (Rl = H).
- L4 ANSWER 7 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Preparation of deuterium labeled styrenes and divinylbenzenes
- AN 1986:515433 CAPLUS
- DN 105:115433
- TI Preparation of deuterium labeled styrenes and divinylbenzenes
- AU Werstiuk, Nick Henry; Timmins, George
- CS Dep. Chem., McMaster Univ., Hamilton, ON, L8S 4M1, Can.
- SO Canadian Journal of Chemistry (1986), 64(6), 1072-6 CODEN: CJCHAG; ISSN: 0008-4042
- DT Journal
- LA English
- OS CASREACT 105:115433
- AB Specifically deuterated styrenes (1-d, 2,2'-d2, and ring-labeled), perdeuterostyrene [19361-62-7], and specifically deuterated divinylbenzenes (1,1'-d2, 2,2,2',2'-d4, and ring-labeled) were prepared by transforming suitably labeled phenylacetic (hydride or deuteride reduction and dehydration by solid KOH) and phenylenediacetic acids (esterification, hydride or deuteride reduction, and dehydration by solid KOH), resp.
- L4 ANSWER 8 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Double homologation of terephthalaldehyde by Wittig and Horner-Wittig reactions: synthesis of 1,4-benzenediacetaldehyde
- AN 1986:497101 CAPLUS
- DN 105:97101
- TI Double homologation of terephthalaldehyde by Wittig and Horner-Wittig reactions: synthesis of 1,4-benzenediacetaldehyde
- AU Campa, Carme; Sanchez-Ferrando, Francisco; Tristan-Polo, Manuel
- CS Fac. Cienc., Univ. Auton. Barcelona, Barcelona, Spain
- SO Nouveau Journal de Chimie (1985), 9(7), 493-8 CODEN: NJCHD4; ISSN: 0398-9836
- DT Journal
- LA English
- OS CASREACT 105:97101
- AB The Wittig reaction of 1,4-C6H4(CHO)2 with equimolar Ph3P:CHOMe (I) orPh2P(O)CH2OMe (II) gave 4-OHCC6H4CH:CHOMe (III), whereas with excess I or II 1,4-C6H4(CH:CHOMe)2 was obtained. Hydrolysis of III gave 4-OCHC6H4CH2CHO. Similarly PhCHO and I gave PhCH:CHOMe, which was hydrolyzed to PhCH2CHO. 1,4-C6H4(CH2CHO)2 was also obtained in 50% yield by reduction of 1,4-C6H4(CH2CO2Et)2 by Dibal.
- L4 ANSWER 9 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Arylacetates
- AN 1983:106993 CAPLUS
- DN 98:106993
- TI Arylacetates
- PA Denki Kagaku Kogyo K. K., Japan
- SO Jpn. Kokai Tokkyo Koho, 5 pp. CODEN: JKXXAF
- DT Patent
- LA Japanese
- FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	JP 57183740	A2	19821112	JP 1982-68850	19820426
	JP 59021852	В4	19840522		
				JP 1982-68850	19820426

AB RCHR1CO2R2 [I, R = (un)substituted aryl, R1 = H, alkyl, R2 = alkyl] were prepared by alkoxycarbonylation of RCHR1R3 (R3 = halo). Thus, stirring PhCH2Cl with MeOH and Na2CO3 in the presence of Co(CO)4- at CO 5 kg/cm2 and 55° for 4 h gave 91% I (R = Ph, R1 = H, R2 = Me).

- L4 ANSWER 10 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Ammonolysis of phenylenediacetic acid esters
- AN 1973:83990 CAPLUS
- DN 78:83990
- TI Ammonolysis of phenylenediacetic acid esters
- AU Ioffe, A. E.; Khcheyan, Kh. E.
- CS Nauchno-Issled. Inst. Sint. Spirtov. Org. Prod., Moscow, USSR
- SO Neftekhimiya (1972), 12(6), 883-93 CODEN: NEFTAH; ISSN: 0028-2421
- DT Journal
- LA Russian
- AB Esterification of m- and p-(HO2CCH2)2C6H4 by excess ROH (R = C1-C10 n-alkyl) in the presence of H2SO4 yielded the corresponding (RO2CCH2)2C6H4 (I) in 89.1-99.0% yield. Ammonolysis of I using 26% aqueous NH3 at 0-5° yielded m- and p-(H2NCOCH2)2C6H4 and m- and p-(NH4+-O2CCH2)2C6H4; the amide-ammonium salt ratio decreased with increasing length of R and temperature at 0-30°.
- L4 ANSWER 11 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Synthesis of esters of phenylenediacetic acids based on diethylbenzene
- AN 1972:153294 CAPLUS
- DN 76:153294
- TI Synthesis of esters of phenylenediacetic acids based on diethylbenzene
- AU Khcheyan, Kh. E.; Ioffe, A. E.; Kostyuk, A. G.
- CS USSR
- SO Khimicheskaya Promyshlennost (Moscow, Russian Federation) (1972), 48(2), 98-100
 CODEN: KPRMAW; ISSN: 0023-110X
- DT Journal
- LA Russian
- GI For diagram(s), see printed CA Issue.
- AB Ten title esters (I, R = n-C1-5 alkyl), having low viscosities and high b.ps., were prepared from the corresponding C6H4Et2 by liquid phase oxidation
- C6H4(COMe)2 with atmospheric O, Willgerodt or Willgerodt-Kindler reaction, and esterification with an alc. The esters were soluble in CHCl3, CCl4, and alcs., but insol. in water.

=> d 14 1-4 ti fbib abs

- L4 ANSWER 1 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Binding of Tetramethylammonium to Polyether Side-Chained Aromatic Hosts. Evaluation of the Binding Contribution from Ether Oxygen Donors
- AN 2003:738628 CAPLUS
- DN 139:364597
- TI Binding of Tetramethylammonium to Polyether Side-Chained Aromatic Hosts. Evaluation of the Binding Contribution from Ether Oxygen Donors
- AU Bartoli, Sandra; De Nicola, Gina; Roelens, Stefano
- CS CNR, Istituto di Chimica dei Composti Organometallici, Dipartimento di Chimica Organica, Universita di Firenze, Sesto Fiorentino, I-50019, Italy
- SO Journal of Organic Chemistry (2003), 68(21), 8149-8156 CODEN: JOCEAH; ISSN: 0022-3263

- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 139:364597
- A set of macrocyclic and open-chain aromatic ligands endowed with polyether AΒ side chains was prepared to assess the contribution of ether O donors to the binding of tetramethylammonium (TMA), a cation believed incapable of interacting with O donors. The open-chain hosts consisted of an aromatic binding site and side chains possessing a variable number of ether O donors; the macrocyclic ligands were based on the structure of a previously studied host, the dimeric cyclophane 1,4-xylylene-1,4-phenylene diacetate (DXPDA), implemented with polyether-type side chains in the backbone. Association to tetramethylammonium picrate (TMAP) was measured in CDC13 at T =296 K by 1H NMR titrns. The main contribution to the binding of TMA comes from the cation- π interaction established with the aromatic binding sites, but they unequivocally show that polyether chains participate with cooperative contributions, although of markedly smaller entity. Water is also bound, but the two quests interact with aromatic rings and O donors in an essentially noncompetitive way. An improved procedure for the preparation of cyclophanic tetraester derivs. was developed that conveniently recycles the oligomeric ester byproducts formed in the 1-pot cyclization reaction. An alternative entry to benzylic diketones also was provided that makes use of a low-order cyanocuprate reagent to prepare in fair yields a class of compds. otherwise uneasily accessible.
- RE.CNT 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L4 ANSWER 2 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Enzymic synthesis of aromatic polyesters by lipase-catalyzed polymerization of dicarboxylic acid divinyl esters and glycols
- AN 1999:258643 CAPLUS
- DN 131:45147
- TI Enzymic synthesis of aromatic polyesters by lipase-catalyzed polymerization of dicarboxylic acid divinyl esters and glycols
- AU Uyama, Hiroshi; Yaguchi, Shigeru; Kobayashi, Shiro
- CS Department of Materials Chemistry, Graduate School of Engineering, Kyoto University, Kyoto, 606-8501, Japan
- SO Polymer Journal (Tokyo) (1999), 31(4), 380-383 CODEN: POLJB8; ISSN: 0032-3896
- PB Society of Polymer Science, Japan
- DT Journal
- LA English
- AB Polyesters containing aromatic moieties in the main chain have been synthesized by enzymic polycondensation (loss of acetaldehyde) of dicarboxylic acid divinyl esters with glycols under mild reaction conditions. Divinyl esters of isophthalic acid, terephthalic acid, and p-phenylenediacetic acid, and p-phenylenedimethanol were used as aromatic monomers. Effects of the polymerization conditions were systematically investigated in the polymerization of
 - divinyl isophthalate and 1,6-hexanediol. Candida antarctica lipase afforded the polymer of the highest mol. weight Methylene chain length of the glycol affected the polymer yield and mol. weight Divinyl terephthalate was enzymically polymerized under similar reaction conditions, yielding polymers of lower mol. weight
- RE.CNT 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L4 ANSWER 3 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Binding of Acetylcholine and Quaternary Ammonium Cations to Macrocyclic and Acyclic "Phane" Esters. Evaluation of the Cation- π Primary Interaction through Adaptive Aromatic Hosts
- AN 1998:739908 CAPLUS
- DN 130:66141

- TI Binding of Acetylcholine and Quaternary Ammonium Cations to Macrocyclic and Acyclic "Phane" Esters. Evaluation of the Cation- π Primary Interaction through Adaptive Aromatic Hosts
- AU Roelens, Stefano; Torriti, Riccardo
- CS Centro di Studio sulla Chimica e la Struttura dei Composti Eterociclici e loro Applicazioni Dipartimento di Chimica Organica, Universita di Firenze, Firenze Italy, I-50121, Italy
- SO Journal of the American Chemical Society (1998), 120(48), 12443-12452 CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- A family of adaptive macrocyclic and acyclic "phane" esters has been AB designed to systematically investigate the interaction between aromatic rings and quaternary ammonium cations in the absence of superimposed contributions, such as hydrophobic, ion-pairing, macrocyclic, and preorganization contributions, to quant. evaluate the primary force at the origin of the cation- π interaction. The unprecedented association with open-chain and cyclic nonpreorganized aromatic hosts in solution is reported, including the remarkable case of binding to phenylacetate ester, that allowed the direct evaluation of the interaction with a single Ph ring. The magnitude of the cation- π attraction has been measured in CDC13 at T = 296 K for picrate salts of acetylcholine (ACh) and tetramethylammonium (TMA), the latter showing the strongest interaction with cyclophane 1b (8.3 kJ mol-1). Results unambiguously confirmed that the basic driving force is a purely electrostatic attraction between the permanent charge of the cation and the aromatic ring. Exptl. standard binding free energies suggest

that interactions of Ph rings are additive, each contributing 2 kJ mol-1 to the overall binding energy, up to a saturation limit in the range of 8 kJ mol-1, consistent with tetracoordinative capabilities of quaternary ammonium cations. Cooperative effects are displayed by the ester group, itself incapable of binding. The possible origin of the ester cooperativity is discussed.

- RE.CNT 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L4 ANSWER 4 OF 11 CAPLUS COPYRIGHT 2004 ACS on STN
- TI Vapor deposition polymerization of dispiro[2.2.2.2]deca-4,9-diene
- AN 1993:449986 CAPLUS
- DN 119:49986
- TI Vapor deposition polymerization of dispiro[2.2.2.2]deca-4,9-diene
- AU Iwatsuki, Shouji; Kubo, Masataka; Hori, Yasutoshi
- CS Fac. Eng., Mie Univ., Tsu, 514, Japan
- SO Macromolecules (1993), 26(6), 1407-10 CODEN: MAMOBX; ISSN: 0024-9297
- DT Journal.
- LA English
- Dispiro[2.2.2.2]deca-4,9-diene (I) was sublimed under a pressure of 0.1 mmHg and was pyrolyzed at 500°. When condensed on a glass surface at 20°, the pyrolyzed gas underwent spontaneous polymerization to give poly(1,4-phenylene-1,2-dimethylethylene) (II) as a film (.hivin.Mn = 3 + 104) in 10-20% yield and oligo(1,4-phenylenetetramethylene-co-1,4-phenylene-1,2-dimethylethylene) as an oil (.hivin.Mn = (2-4) + 102) in 50-70% yield. It was proposed for the formation of the polymer film that the diradical intermediate generated in the pyrolysis of I underwent an isomerization reaction to form 7,8-dimethyl-1,4-benzoquinodimethane which polymerized spontaneously to give II film.

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NEWS 7 MAR 02 GBFULL: New full-text patent database on STN
NEWS 8 MAR 03 REGISTRY/ZREGISTRY - Sequence annotations enhanced NEWS 9 MAR 03 MEDLINE file segment of TOXCENTER reloaded NEWS 10 MAR 22 KOREAPAT now updated monthly; patent information enhanced NEWS 11 MAR 22 Original IDE display format returns to REGISTRY/ZREGISTRY NEWS 12 MAR 22 PATDPASPC - New patent database available NEWS 13 MAR 22 REGISTRY/ZREGISTRY enhanced with experimental property tags NEWS 14 APR 04 EPFULL enhanced with additional patent information and new NEWS 15 APR 04 EMBASE - Database reloaded and enhanced NEWS 16 APR 18 New CAS Information Use Policies available online

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NEWS 2 "Ask CAS" for self-help around the clock

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NEWS 4 FEB 28 PATDPAFULL - New display fields provide for legal status data from INPADOC

NEWS 5 FEB 28 BABS - Current-awareness alerts (SDIs) available

NEWS 6 FEB 28 MEDLINE/LMEDLINE reloaded

NEWS 7 MAR 02 GBFULL: New full-text patent database on STN

NEWS 8 MAR 03 REGISTRY/ZREGISTRY - Sequence annotations enhanced

NEWS 9 MAR 03 MEDLINE file segment of TOXCENTER reloaded

NEWS 10 MAR 22 KOREAPAT now updated monthly; patent information enhanced

NEWS 11 MAR 22 Original IDE display format returns to REGISTRY/ZREGISTRY

NEWS 12 MAR 22 PATDPASPC - New patent database available

NEWS 13 MAR 22 REGISTRY/ZREGISTRY enhanced with experimental property tags

NEWS 14 APR 04 EPFULL enhanced with additional patent information and new fields

NEWS 15 APR 04 EMBASE - Database reloaded and enhanced

NEWS 16 APR 18 New CAS Information Use Policies available online

NEWS 17 APR 25 Patent searching, including current-awareness alerts (SDIs),

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=> phenyalanine

L1 108 PHENYALANINE

=> phenylalanine

73297 PHENYLALANINE

733 PHENYLALANINES

L2 73592 PHENYLALANINE

(PHENYLALANINE OR PHENYLALANINES)

=> diabet?

L3 111423 DIABET?

=> 12 and 13

L4 553 L2 AND L3

=> d 14 543-553 ti

L4 ANSWER 543 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN

TI Amino acid metabolism studies with the isolated perfused rat liver

L4 ANSWER 544 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN .

TI Amino acid metabolism in diabetes mellitus

- L4 ANSWER 545 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Incorporation of Cl4-amino acids into glutathione and protein fractions of normal and diabetic rat tissues
- L4 ANSWER 546 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Amino acid metabolism in disease of the liver
- L4 ANSWER 547 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Diabetogenic activity as an inherent property of growth hormone
- L4 ANSWER 548 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Free plasma levels and urinary excretion of eighteen amino acids in normal and diabetic dogs
- L4 ANSWER 549 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Excretion of free amino acids in alloxan diabetes
- L4 ANSWER 550 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Influence of protein on the ketone body elimination in severe diabetes
- L4 ANSWER 551 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI "Acetone bodies" in diabetes mellitus
- L4 ANSWER 552 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI The Breaking Down of Fatty Acids in Diabetes Mellitus. IV
- L4 ANSWER 553 OF 553 CAPLUS COPYRIGHT 2005 ACS on STN
- TI The Catabolism of Fatty Acids in Diabetes Mellitus. II
- => logofdf hold

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34551 HOLD

24055 HOLDS

57735 HOLD

(HOLD OR HOLDS)

L5 0 LOGOFDF HOLD

(LOGOFDF(W)HOLD)

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SINCE FILE

TOTAL

FULL ESTIMATED COST

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FULL ESTIMATED COST	15.22	15.43

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chain nodes :
7 8 9 10 11 12 13 16
ring nodes :
1 2 3 4 5 6
chain bonds :
1-9 4-7 7-8 8-11 9-10 9-16 10-12 10-13
ring bonds :
1-2 1-6 2-3 3-4 4-5 5-6
exact/norm bonds :
8-11 9-16 10-12 10-13
exact bonds :
1-9 4-7 7-8 9-10
normalized bonds :
1-2 1-6 2-3 3-4 4-5 5-6

Match level:
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS 12:CLASS 13:CLASS 16:CLASS

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L6 HAS NO ANSWERS
L6 STR

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1: ANSWERS

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75.1% PROCESSED 1000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

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FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 24432 TO 28808 PROJECTED ANSWERS: 63 TO 521

L7 11 SEA SSS SAM L6

=> d scan

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN 1,4-Benzenediacetic acid, α -amino-, (S)- (9CI)

MF C10 H11 N O4

Absolute stereochemistry. Rotation (+).

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):11

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN 1,4-Benzenediacetic acid, α -(benzoylamino)-, dimethyl ester (9CI)

MF C19 H19 N O5

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, α -[bis(phenylmethyl)amino]-4-carboxy- (9CI)

MF C23 H21 N O4

CI COM

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, $\alpha-[(1,1-dimethylethoxy)carbonyl]amino]-4-(methoxycarbonyl)- (9CI)$

MF C15 H19 N O6

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, α -[(aminocarbonyl)amino]-4-(methoxycarbonyl)-,

(R)- (9CI)

MF C11 H12 N2 O5

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT.

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN 1,4-Benzenediacetic acid, α,α' -bis[bis[2-(1,3,2-benzodioxazol-

2-yl)ethyl]amino]-, diethyl ester (9CI)

MF C46 H48 N6 O12

PAGE 1-B

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L7 11.ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, α -amino-4-carboxy- α -ethyl- (9CI)

MF C11 H13 N O4

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, 4-formyl-3-hydroxy-α-[[[(4methoxyphenyl)methoxy]carbonyl]amino]-, compd. with Ncyclohexylcyclohexanamine (1:1) (9CI)

MF C18 H17 N O7 . C12 H23 N

CM 1

CM 2

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, 4-(aminocarbonyl)- α -[[(1,1-dimethylethoxy)carbonyl]amino]-, (α R)- (9CI) MF C14 H18 N2 O5

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN 1H-Indole-4-acetic acid, α -amino-7-carboxy- (9CI)

MF C11 H10 N2 O4

L7 11 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN 1,4-Benzenediacetic acid, α,α' -bis[[(1,1-dimethylethoxy)carbonyl]amino]-, [S-(R*,R*)]- (9CI) MF C20 H28 N2 O8

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

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SINCE FILE TOTAL ENTRY SESSION 0.86 16.29

FULL ESTIMATED COST

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FILE COVERS 1907 - 26 Apr 2005 VOL 142 ISS 18 FILE LAST UPDATED: 25 Apr 2005 (20050425/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 17

L8 18 L7

=> d his

(FILE 'HOME' ENTERED AT 10:44:00 ON 26 APR 2005)

FILE 'CAPLUS' ENTERED AT 10:44:33 ON 26 APR 2005

L1 108 PHENYALANINE

L2 73592 PHENYLALANINE

L3 111423 DIABET?

L4 553 L2 AND L3

L5 0 LOGOFDF HOLD

FILE 'REGISTRY' ENTERED AT 10:53:30 ON 26 APR 2005

L6 STRUCTURE UPLOADED

L7 11 SEARCH L6 SSS SAM

FILE 'CAPLUS' ENTERED AT 10:54:54 ON 26 APR 2005

L8 18 L7

=> 13 and 18

L9 0 L3 AND L8

=> file req

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION

16.74

0.45

FULL ESTIMATED COST

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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0 DICTIONARY FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

*

* The CA roles and document type information have been removed from * the IDE default display format and the ED field has been added, * effective March 20, 2005. A new display format, IDERL, is now * available and contains the CA role and document type information. *

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> search 16 sss full FULL SEARCH INITIATED 10:57:13 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 26327 TO ITERATE

100.0% PROCESSED 26327 ITERATIONS SEARCH TIME: 00.00.01

228 ANSWERS

L10 228 SEA SSS FUL L6

=> d scan

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, α -amino-4-carboxy-2-[methyl(phenylmethyl)amino]-(9CI)

MF C17 H18 N2 O4

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):20

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, $\alpha-[[(1,1-\text{dimethylethoxy})\,\text{carbonyl}]\,\text{amino}]-4-(\text{methoxycarbonyl})-, \text{methyl ester (9CI)}$

MF C16 H21 N O6

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, $\alpha\text{-amino-4-(methoxycarbonyl)-, perchlorate (9CI)}$ MF C10 H11 N O4 . C1 H O4

CM 1

CM 2

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -[(diphenylmethylene)amino]-4-(methoxycarbonyl)-, ethyl ester (9CI) MF C25 H23 N O4

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -methyl-4-[(1-methylethoxy)carbonyl]- α -[(trifluoroacetyl)amino]-, 1-methylethyl ester (9CI) MF C18 H22 F3 N O5

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -amino-4-carboxy-, (α S)- (9CI) MF C9 H9 N O4

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -amino-4-formyl-3-hydroxy-, (α S)- (9CI) MF C9 H9 N O4

Absolute stereochemistry.

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, $\alpha-[[(1,1-\text{dimethylethoxy})\text{carbonyl}]\text{amino}]-4-(methoxycarbonyl)-, (<math>\alpha R$)- (9CI)

MF C15 H19 N O6

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN 1,4-Benzenediacetic acid, α -[[2-fluoro-4-[methyl(4,6,7,8-tetrahydro-

2-methyl-4-oxo-1H-cyclopenta[g]quinazolin-6-yl)amino]benzoyl]amino]- (9CI)

MF C30 H27 F N4 O6

PAGE 1-A

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

Benzeneacetic acid, α -[(chloroacetyl)amino]-4-formyl-3-hydroxy-, IN compd. with N-cyclohexylcyclohexanamine (1:1) (9CI) C12 H23 N . C11 H10 C1 N O5

MF

CM 1

CM 2

REGISTRY COPYRIGHT 2005 ACS on STN L10 228 ANSWERS

IN 1,4-Benzenediacetic acid, α,α' -diamino-, diethyl ester (9CI)

MF C14 H20 N2 O4

COM CI

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -amino-4-carboxy-2-methyl-, (R)- (9CI)

MF C10 H11 N O4

Absolute stereochemistry. Rotation (-).

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, α -amino-4-carboxy-3-hydroxy- α -methyl-(9CI)

MF C10 H11 N O5

$$\begin{array}{c|c} & \text{NH2} \\ & \text{C-CO}_2\text{H} \\ \\ \text{HO}_2\text{C} & \text{OH} \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN 1,4-Benzenediacetic acid, α,α' -bis[(4-bromophenyl)amino]-(9CI)

MF C22 H18 Br2 N2 O4

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Nitroxide, carboxy(4-carboxyphenyl)methyl 1,1-dimethyl-2-(octylthio)ethyl,

ion(1-) (9CI) MF C21 H31 N O5 S CI COM

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -amino-4-carboxy-2-phenoxy- (9CI) MF C15 H13 N O5

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -amino-4-(methoxycarbonyl)-, methyl ester, (R)-, perchlorate (9CI) MF C11 H13 N O4 . Cl H O4

CM 1

Absolute stereochemistry.

CM 2

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

Benzeneacetic acid, 4-carboxy-2-methoxy- α -[(phenylacetyl)amino]-

(9CI)

C18 H17 N O6 MF

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L10 REGISTRY COPYRIGHT 2005 ACS on STN 228 ANSWERS

IN Benzeneacetic acid, 4-carboxy- α -[[(1,1-dimethylethoxy)carbonyl]amino

]-, α -methyl ester (9CI)

MF C15 H19 N O6

L10 228 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN IN Benzeneacetic acid, α -amino-4-carboxy-3-hydroxy- (9CI) MF C9 H9 N O5

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):file caplus 'FILE CAPLUS' IS NOT VALID HERE

To display more answers, enter the number of answers you would like to see. To end the display, enter "NONE", "N", "0", or "END". HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> file caplus COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
162.19 178.93

FULL ESTIMATED COST

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FILE COVERS 1907 - 26 Apr 2005 VOL 142 ISS 18 FILE LAST UPDATED: 25 Apr 2005 (20050425/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 110 L11

337 L10

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(FILE 'HOME' ENTERED AT 10:44:00 ON 26 APR 2005)
     FILE 'CAPLUS' ENTERED AT 10:44:33 ON 26 APR 2005
L1
           108 PHENYALANINE
         73592 PHENYLALANINE
L2
         111423 DIABET?
L3
            553 L2 AND L3
L4
             0 LOGOFDF HOLD
L5
     FILE 'REGISTRY' ENTERED AT 10:53:30 ON 26 APR 2005
                STRUCTURE UPLOADED
L6
L7
             11 SEARCH L6 SSS SAM
     FILE 'CAPLUS' ENTERED AT 10:54:54 ON 26 APR 2005
L8
            18 L7
             0 L3 AND L8
L9
     FILE 'REGISTRY' ENTERED AT 10:55:45 ON 26 APR 2005
           228 SEARCH L6 SSS FULL
L10
     FILE 'CAPLUS' ENTERED AT 10:57:43 ON 26 APR 2005
L11
           337 L10
=> 13 and 111
           0 L3 AND L11
=> phenyl acetic
        321396 PHENYL
           384 PHENYLS
        321656 PHENYL
                (PHENYL OR PHENYLS)
      1225166 PH
         9328 PHS
       1229262 PH
                 (PH OR PHS)
       1465011 PHENYL
                 (PHENYL OR PH)
        211465 ACETIC
            22 ACETICS
        211474 ACETIC
                 (ACETIC OR ACETICS)
L13
           875 PHENYL ACETIC
                 (PHENYL (W) ACETIC)
=> phenylacetic
L14
      .8713 PHENYLACETIC
=> d hid
'HID' IS NOT A VALID FORMAT FOR FILE 'CAPLUS'
The following are valid formats:
ABS ----- GI and AB
ALL ----- BIB, AB, IND, RE
APPS ----- AI, PRAI
BIB ----- AN, plus Bibliographic Data and PI table (default)
CAN ----- List of CA abstract numbers without answer numbers
CBIB ----- AN, plus Compressed Bibliographic Data
DALL ----- ALL, delimited (end of each field identified)
DMAX ----- MAX, delimited for post-processing
```

FAM ----- AN, PI and PRAI in table, plus Patent Family data

```
FBIB ----- AN, BIB, plus Patent FAM
IND ----- Indexing data
IPC ----- International Patent Classifications
MAX ----- ALL, plus Patent FAM, RE
PATS ----- PI, SO
SAM ----- CC, SX, TI, ST, IT
SCAN ----- CC, SX, TI, ST, IT (random display, no answer numbers;
             SCAN must be entered on the same line as the DISPLAY,
             e.g., D SCAN or DISPLAY SCAN)
STD ---- BIB, IPC, and NCL
IABS ----- ABS, indented with text labels
IALL ----- ALL, indented with text labels
IBIB ----- BIB, indented with text labels
IMAX ----- MAX, indented with text labels
ISTD ----- STD, indented with text labels
OBIB ----- AN, plus Bibliographic Data (original)
OIBIB ----- OBIB, indented with text labels
SBIB ----- BIB, no citations
SIBIB ----- IBIB, no citations
HIT ----- Fields containing hit terms
HITIND ----- IC, ICA, ICI, NCL, CC and index field (ST and IT)
             containing hit terms
HITRN ----- HIT RN and its text modification
HITSTR ----- HIT RN, its text modification, its CA index name, and
             its structure diagram
HITSEQ ----- HIT RN, its text modification, its CA index name, its
             structure diagram, plus NTE and SEQ fields
FHITSTR ---- First HIT RN, its text modification, its CA index name, and
             its structure diagram
FHITSEQ ---- First HIT RN, its text modification, its CA index name, its
             structure diagram, plus NTE and SEQ fields
KWIC ----- Hit term plus 20 words on either side
OCC ----- Number of occurrence of hit term and field in which it occurs
```

To display a particular field or fields, enter the display field codes. For a list of the display field codes, enter HELP DFIELDS at an arrow prompt (=>). Examples of formats include: TI; TI,AU; BIB,ST; TI,IND; TI,SO. You may specify the format fields in any order and the information will be displayed in the same order as the format specification.

All of the formats (except for SAM, SCAN, HIT, HITIND, HITRN, HITSTR, FHITSTR, HITSEQ, FHITSEQ, KWIC, and OCC) may be used with DISPLAY ACC to view a specified Accession Number. ENTER DISPLAY FORMAT (BIB):end

=> d his

L6

(FILE 'HOME' ENTERED AT 10:44:00 ON 26 APR 2005)

FILE 'REGISTRY' ENTERED AT 10:53:30 ON 26 APR 2005 STRUCTURE UPLOADED

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11 SEARCH L6 SSS SAM
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FILE 'CAPLUS' ENTERED AT 10:54:54 ON 26 APR 2005 18 L7

L9 0 L3 AND L8

FILE 'REGISTRY' ENTERED AT 10:55:45 ON 26 APR 2005 L10 228 SEARCH L6 SSS FULL

FILE 'CAPLUS' ENTERED AT 10:57:43 ON 26 APR 2005

L11 337 L10

L12 0 L3 AND L11 L13 875 PHENYL ACETIC L14 8713 PHENYLACETIC

=> 13 and 114

L7

 $rac{1}{8}$

L15 107 L3 AND L14

=> d 97-107 til15

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The following are valid formats:

ABS ----- GI and AB

ALL ----- BIB, AB, IND, RE

APPS ----- AI, PRAI

BIB ----- AN, plus Bibliographic Data and PI table (default)

CAN ----- List of CA abstract numbers without answer numbers

CBIB ----- AN, plus Compressed Bibliographic Data

DALL ----- ALL, delimited (end of each field identified)

DMAX ----- MAX, delimited for post-processing

FAM ----- AN, PI and PRAI in table, plus Patent Family data

FBIB ----- AN, BIB, plus Patent FAM

IND ----- Indexing data

IPC ----- International Patent Classifications

MAX ----- ALL, plus Patent FAM, RE

PATS ----- PI, SO

SAM ----- CC, SX, TI, ST, IT

SCAN ----- CC, SX, TI, ST, IT (random display, no answer numbers; SCAN must be entered on the same line as the DISPLAY,

e.g., D SCAN or DISPLAY SCAN)

STD ----- BIB, IPC, and NCL

IABS ----- ABS, indented with text labels

IALL ----- ALL, indented with text labels

IBIB ----- BIB, indented with text labels

IMAX ----- MAX, indented with text labels

ISTD ----- STD, indented with text labels

OBIB ----- AN, plus Bibliographic Data (original)

OIBIB ----- OBIB, indented with text labels

SBIB ----- BIB, no citations

SIBIB ----- IBIB, no citations

HIT ----- Fields containing hit terms

HITIND ----- IC, ICA, ICI, NCL, CC and index field (ST and IT)

containing hit terms

HITRN ----- HIT RN and its text modification

HITSTR ----- HIT RN, its text modification, its CA index name, and

its structure diagram

HITSEQ ----- HIT RN, its text modification, its CA index name, its structure diagram, plus NTE and SEQ fields

FHITSTR ---- First HIT RN, its text modification, its CA index name, and its structure diagram

FHITSEQ ---- First HIT RN, its text modification, its CA index name, its structure diagram, plus NTE and SEQ fields

KWIC ----- Hit term plus 20 words on either side

OCC ----- Number of occurrence of hit term and field in which it occurs

To display a particular field or fields, enter the display field codes. For a list of the display field codes, enter HELP DFIELDS at an arrow prompt (=>). Examples of formats include: TI; TI, AU; BIB, ST; TI, IND; TI, SO. You may specify the format fields in any order and the information will be displayed in the same order as the format specification.

All of the formats (except for SAM, SCAN, HIT, HITIND, HITRN, HITSTR, FHITSTR, HITSEQ, FHITSEQ, KWIC, and OCC) may be used with DISPLAY ACC to view a specified Accession Number.
ENTER DISPLAY FORMAT (BIB):end

=> d 115 97-107 ti

- L15 ANSWER 97 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Benzothiazol-2-ylcarboxylic acids with diverse spacers: a novel class of potent, orally active aldose reductase inhibitors
- L15 ANSWER 98 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI In vitro and in vivo suppression of gluconeogenesis by inhibition of pyruvate carboxylase
- L15 ANSWER 99 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI 4-[2-(2-hydroxy-2-phenylethylamino)ethyl]**phenylacetic** acid as β3-adrenoceptor agonist
- L15 ANSWER 100 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Compositions and methods for treating autoimmune diseases
- L15 ANSWER 101 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Cyclohexylamides and tachykinin inhibitors containing the cyclohexylamides for pharmaceutical preparations
- L15 ANSWER 102 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of N-substituted heterocyclic derivatives and their pharmaceutical compositions as angiotensin II receptor antagonists
- L15 ANSWER 103 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of (phenylalkyl)propanolamine derivatives as antidiabetics as antiobesity agents
- L15 ANSWER 104 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Urinary organic acids in natural early-onset insulin-dependent diabetic dogs
- L15 ANSWER 105 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI N-benzyl-2-phenylacetamide derivatives and their use as hypoglycemics
- L15 ANSWER 106 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Pharmacological studies on oral hypoglycemic agents
- L15 ANSWER 107 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Proteins and the Deposition of Fat in the Liver

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ANSWER 99 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
     4-[2-(2-hydroxy-2-phenylethylamino)ethyl]phenylacetic acid as
TI
     β3-adrenoceptor agonist
AN
     1994:244357 CAPLUS
DN
     120:244357
     4-[2-(2-hydroxy-2-phenylethylamino)ethyl]phenylacetic acid as
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     β3-adrenoceptor agonist
IN
     Holloway, Brian Roy; Howe, Ralph; Rao, Balbir Singh
PA
     Zeneca Ltd., UK
SO
     PCT Int. Appl., 44 pp.
     CODEN: PIXXD2
ÐΤ
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FAN.CNT 1
     PATENT NO.
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                                 DATE
                                              APPLICATION NO.
                                                                      DATE
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                                 19931111
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                          A1
                                                                      19930420
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                                              GB 1992-9076
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                                                                      19920427
                                                                  Α
                                              WO 1993-GB821
                                                                   W
                                                                      19930420
     CASREACT 120:244357; MARPAT 120:244357
os
GI
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Phch (OH)
$$CH_2NHCH_2CH_2$$
 — CH_2CO_2H

- AB The title compound I and in-vivo hydrolyzable esters and pharmaceutically acceptable salts are prepared and shown to have β 3-adrenoceptor agonist activity and antiobesity, hypoglycemic, and related therapeutic utilities. I intermediates are also prepared and I-containing formulations presented.
- L15 ANSWER 103 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of (phenylalkyl)propanolamine derivatives as antidiabetics as antiobesity agents
- AN 1990:234956 CAPLUS
- DN 112:234956
- TI Preparation of (phenylalkyl)propanolamine derivatives as antidiabetics as antiobesity agents
- IN Kienzle, Frank

PA Hoffmann-La Roche, F., und Co. A.-G., Switz.

SO Eur. Pat. Appl., 14 pp.

CODEN: EPXXDW

DT Patent LA German

FAN CNT 1

FAIN.	CNT PA	TENT NO.		KIND	DATE	APPLICATION NO.	DATE
PI		345591 345591		A1 B1	19891213 19930331	EP 1989-109675	19890530
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		•		·			19880610
	FI	8902341		Α	19891211	FI 1989-2341	19890516
						CH 1988-2245 F	19880610
	AT	87610		E	19930415	AT 1989-109675	19890530
						CH 1988-2245	19880610
						EP 1989-109675 A	19890530
	ES	2053866		Т3	19940801	ES 1989-109675	19890530
						· CH 1988-2245	19880610
	ZA	8904210		Α	19900328	ZA 1989-4210	19890602
						CH 1988-2245 A	19880610
	AU	8936026		A1	19891214	AU 1989-36026	19890605
	ΑU	622907		B2	19920430		
						CH 1988-2245	19880610
	HU	55344		A2	19910528	HU 1989-2868	19890605
							19880610
	JP	02036158		A2	19900206	JP 1989-144282	19890608
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	DK	8902842		Α	19891211	DK 1989-2842	19890609
			-	•			19880610
		8902387		Α	19891211	NO 1989-2387	19890609
		170011		В	19920525		
	МО	170011		С	19920902		
							19880610
	US	5045567	•	A	19910903	US 1990-608610	19901031
							19880610
						US 1989-363242 F	81 19890608 🍻

OS MARPAT 112:234956

GI

$$\text{HOCH}_2\text{CHR}^2\text{CH}_2\text{NR}^1\text{CHR}^3\text{CH}_2$$
 \longrightarrow OR^4

The title compds. I [R1 = H or CH2CHR5(CH2)nOH, R5 = Ph, m-halophenyl, m-F3CC6H4, thienyl, or pyridyl; R2 = R5; R3 = H, Me; R4 = H, HO2CCH2, C1-4 alkoxycarbonylmethyl, C1-4 alkoxyethyl, or Ph C1-4 alkyloxyethyl) and their compatible physiol. salts having a catabolic effect are prepared for use in the treatment of obesity, diabetes mellitus, conditions involving increased protein degradation, and as food additives for obese animals. Thus, di-Et phenylmalonate in diglyme was treated with p-(2-ethoxyethoxy)phenethylamine, the solution stirred 48 h at 95°, cooled, the solvent removed, and the residue chromatog. purified to give Et [[[p-(2-ethoxyethoxy)phenethyl]carbamoyl]phenyl]acetate (II). The effects of II on the O consumption of albino rats showed its effectiveness in treating obesity.

L15 ANSWER 104 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN

- TI Urinary organic acids in natural early-onset insulin-dependent diabetic dogs
- AN 1988:627815 CAPLUS
- DN 109:227815
- TI Urinary organic acids in natural early-onset insulin-dependent diabetic dogs
- AU Shigematsu, Yosuke; Sweeley, Charles C.; Schall, William D.; Gossain, Ved
- CS Dep. Biochem., Michigan State Univ., East Lansing, MI, 48824, USA
- SO Acta Paediatrica Japonica (1988), 30(3), 285-93 CODEN: APDJBE; ISSN: 0374-5600
- DT Journal
- LA English
- AB The urinary organic acids of spontaneously-occurring, insulin-dependent diabetic dogs under insulin therapy were compared with those of normal dogs, using a semi-automated sample injection-capillary gas chromatograph-computerized data processing system. The following acids were excreted in significantly greater amts. by diabetic dogs: 2-hydroxybutyric, 4-deoxytetronic, 3-hydroxybutyric, acetoacetic, arabinonic, erythronic, 3-deoxytetraonic, 2-deoxyribonic, lactic, pyruvic, 2-hydroxyisobutyric and 2-hydroxyisovaleric acids. Not only ketone bodies, but also the metabolites of threonine, 2-hydroxybutyric acid and 4-deoxytetronic acids appear to be important and sensitive markers for the metabolic state in insulin-treated diabetic dogs, although the changes in these acids are not always well correlated with each other.
- L15 ANSWER 105 OF 107 CAPLUS COPYRIGHT 2005 ACS on STN
- TI N-benzyl-2-phenylacetamide derivatives and their use as hypoglycemics
- AN 1987:138092 CAPLUS
- DN 106:138092
- TI N-benzyl-2-phenylacetamide derivatives and their use as hypoglycemics
- IN Grell, Wolfgang; Hurnaus, Rudolf; Sauter, Robert; Reiffen, Manfred;
 Rupprecht, Eckhard
- PA Thomae, Dr. Karl, G.m.b.H., Fed. Rep. Ger.
- SO Ger. Offen., 29 pp.

CODEN: GWXXBX

- DT Patent
- LA German
- FAN.CNT 3

	PAT	TENT NO.			KINI	DATE		API	PLICATION NO.		DATE
ΡI	DE	3523466			A1	1987	0108	DE	1985-3523466		19850701
	EP	208200			A1	1987	0114	EP	1986-108640		19860625
	EP	208200			В1		0425				
								T.T. T.I	J, NL, SE		
			, 5-,	٠,	,	 , 02,	,	-	1985-3523466	А	19850701
	ΤΔ	52255			Е	1990	0515		1986-108640	••	19860625
	111	32233			-	1330			1985-3523466	Δ	19850701
									1986-108640	A	19860625
	CA	1320723			A1	1993	0727		1986-512614	А	19860627
	CA	1320723			ΛŢ	1000	0121		1985-3523466	7\	19850701
	שמ	8603109			Α	1987	0102		1986-3109	A	19860630
								אט	1900-3109		19000030
	DΚ	168741			В1	1994	0530		1005 2502466	_	10050701
					_				1985-3523466		
		8602764			Α	1987		FI	1986-2764		19860630
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			•					DE	1985-3523466	Α	19850701
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	NO	167736			В	1991	0826				
	NO	167736			С	1991	1204				
		· · - •			-	-		DE	1985-3523466	А	19850701
	ΔII	8659383			A1	1987	0108		1986-59383		19860630
		587263			B2	1989		710	1500 55505		1300000
	ΔU	201203			22	エンロラ	~ 0.70			•	

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				DE 1985-3523466	Α	19850701
	JP 62005974	A2	19870112	JP 1986-151864		19860630
	JP 07039406	В4	19950501			
				DE 1985-3523466	Α	19850701
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	ни 196193	В	19881028			
	130130	_		DE 1985-3523466	Α	19850701
	ES 2000443	A6	19880301	ES 1986-56		19860630
	L5 2000415	AU	17000301	DE 1985-3523466	Α	19850701
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	Ta 0003600	3.6	10001101		Α	19850701
	ES 2003629	A6	19881101	ES 1986-3480		19861218
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	ES 2003758	A6	19881116	ES 1986-3481		19861218
				DE 1985-3523466	Α	19850701
	US 5216167	Α	19930601	US 1990-495820		19900621
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				US 1984-684054	B2	19841210
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	05 6143769	A	20001107		70	
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				DE 1985-3523466	Α	19850701
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				US 1986-878921		19860626
				US 1989-302022		19890125
				US 1990-495820		19900621
				US 1984-684054		19941210
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LAIN	PATENT NO.	KIND	חאיייבי	ADDITCATION NO		האתב
	PAIENI NO.	 VIND	DATE	APPLICATION NO.		DATE
DT			10050717	DE 1003 3347565	- -	
PI	DE 3347565	A1	19850711			19831230
	DK 8406131	A	19850701	DK 1984-6131		19841220
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	JP 60158171	A2	19850819	JP 1984-272685		19841224
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	EP 147850	В1	19890614			
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ES 539078	A1	19860316	ES 1984-539078 DE 1983-3347565	19841227 A 19831230
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NO 162819	С	19900221		
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ни 37773 ни 194548	A2 B	19860228 19880229	HU 1984-4870	19841228
NU 194348	В	13000223	DE 1983-3347565	A 19831230
ZA 8410103	Α	19860924	ZA 1984-10103	19841228
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E3 34300U	. AI	19000201	DE 1983-3347565	A 19831230
ES 545881	A1	19860201	ES 1985-545881	19850805
			DE 1983-3347565	A 19831230
ES 545882	A1	19860201	ES 1985-545882	19850805 ··
ES 545883	A1	19860201	DE 1983-3347565 ES 1985-545883	A 19831230 19850805
TP 242002	A.	13000201	DE 1983-3347565	A 19831230
US 5216167	Α	19930601	US 1990-495820	19900621
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			US 1986-878921	B2 19860626
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			US 1986-872706	B2 19860610
			US 1986-878921	B2 19860626
			US 1989-302022 US 1990-495820	B1 19890125 A2 19900621
			ひろ エラブローダブプログロ	NC 1330007

FAN	US 37035		E	20010130	US 1997-946602 DE 1983-3347565 DE 1985-3522604 DE 1985-3523466 US 1986-872706 US 1986-878921 US 1989-302022 US 1990-495820	A A A B2 B2 B1 A5	19920724 19971007 19831230 19850625 19850701 19860610 19860626 19890125 19900621 19941210
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	EP 207331 EP 207331		Al Bl	19870107 19900523			19860610
		BE, CH,				Α	19850625
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•	FI 8602650 FI 82689 FI 82689		A B C	19861226 19901231 19910410	FI 1986-2650	Α	19860623
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	JP 62000474 JP 07039405		A2 B4	19870106 19950501		А	19850625 19860624
	AU 8659139		A1	19870108	DE 1985-3522604 AU 1986-59139	Α	19850625 19860624
	AU 583631 ZA 8604695		B2 A	19890504 19880224	DE 1985-3522604 ZA 1986-4695	Α	19850625 19860624
	ES 556495		A1	19880401	DE 1985-3522604 ES 1986-556495	A	19850625 19860624
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	US 6143769		Α	20001107			19940509

DE 1983-3347565 A 19831230 US 1984-684054 B2 19841210 DE 1985-3522604 A 19850625 DE 1985-3523466 19850701 Α US 1986-872706 B2 19860610 US 1986-878921 B2 19860626 US 1989-302022 B1 19890125 US 1990-495820 A2 19900621 US 1992-919820 Al 19920724 US 37035 Ε 20010130 US 1997-946602 19971007 DE 1983-3347565 19831230 DE 1985-3522604 Α 19850625 DE 1985-3523466 19850701 Α US 1986-872706 B2 19860610 US 1986-878921 B2 19860626 US 1989-302022 B1 19890125 US 1990-495820 A5 19900621 US 1984-684054 B2 19941210

GI

AB The title compds. [I; R1 = (un)substituted alkylenimino; R2 = H, Me, MeO, halo; R3 = H, CO2H, alkoxycarbonyl, (un)substituted alkyl, Ph; R4 = H, alkyl, CH2:CHCH2; W = CHO, CO2H, R5CH2, R6CH2CH2, R7CH:CH; R5 = H, OH, CO2H, cyano; R6 = CO2H, cyano; R7 = CO2H, cyano, alkoxycarbonyl] and their enantiomers and salts were prepared as hypoglycemic agents.
α-(Cyclohexylmethyl)-2-piperidinobenzylamine was amidated with 3-(EtO)-4-(EtO2C)C6H3CH2CO2H (68%) and the product was saponified to give 82% title compound II. In rats 0.5 mg II/kg orally reduced blood sugar 22% after 1 h and 45% after 4 h.

=> logoff hold		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	34.44	213.37
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-2.92	-2.92

SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 11:08:25 ON 26 APR 2005

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID: SSSPTA1623PAZ

PASSWORD:

* * * * * RECONNECTED TO STN INTERNATIONAL * * * * * * SESSION RESUMED IN FILE 'CAPLUS' AT 11:32:11 ON 26 APR 2005 FILE 'CAPLUS' ENTERED AT 11:32:11 ON 26 APR 2005 COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	34.44	213.37
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-2.92	-2.92
=> logoff hold		
COST IN U.S. DOLLARS	SINCE FILE	\mathtt{TOTAL}
	ENTRY	SESSION
FULL ESTIMATED COST	34.89	213.82
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	_2.92	-2.92

SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 11:32:40 ON 26 APR 2005

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID:SSSPTA1623PAZ

PASSWORD:

* * * * * RECONNECTED TO STN INTERNATIONAL * * * * * * * SESSION RESUMED IN FILE 'CAPLUS' AT 12:14:48 ON 26 APR 2005 FILE 'CAPLUS' ENTERED AT 12:14:48 ON 26 APR 2005 COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	34.89	213.82
DISCOUNT AMOUNTS (FOR OUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
, , , , , , , , , , , , , , , , , , , ,	ENTRY	SESSION
CA SUBSCRIBER PRICE	-2.92	-2.92
GI DOBBONIDEN TRICE	,2.32	2.32
=> file reg		
COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	34.89	213.82
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-2.92	-2.92

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STRUCTURE FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0 DICTIONARY FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0

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TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

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=>

Uploading C:\Documents and Settings\PZucker\My Documents\Examination Auxillary files\10797458\10797458 modified subgenus 2.str

chain nodes : 2 3 4 5 6 7 14 15 16 21 22 ring nodes : 1 8 9 10 11 12 chain bonds : 1-2 2-3 3-4 4-5 4-14 5-6 6-7 7-15 15-16 16-21 16-22 ring bonds : 1-8 1-12 8-9 9-10 10-11 11-12 exact/norm bonds : 2-3 3-4 4-14 5-6 6-7 16-21 16-22 exact bonds : 1-2 4-5 7-15 15-16 normalized bonds :

1-8 1-12 8-9 9-10 10-11 11-12

G1:C,O,S,N,SO2

Match level :

1:Atom 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:Atom 7:CLASS 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 14:CLASS 15:CLASS 16:CLASS 21:CLASS 22:CLASS

Generic attributes :

Number of Carbon Atoms : less than 7 Type of Ring System : Monocyclic

L16 STRUCTURE UPLOADED

=> d 116L16 HAS NO ANSWERS STR L16

$$G1$$
 Cy
 0
 0
 0
 0

G1 C, O, S, N, SO2

Structure attributes must be viewed using STN Express query preparation.

=> search 116 sss sam SAMPLE SEARCH INITIATED 12:15:21 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 65654 TO ITERATE

1000 ITERATIONS 1.5% PROCESSED INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

2 ANSWERS

FULL FILE PROJECTIONS:

ONLINE **INCOMPLETE** **INCOMPLETE** BATCH

PROJECTED ITERATIONS:

EXCEEDS 1000000

PROJECTED ANSWERS:

EXCEEDS 1939

L17

2 SEA SSS SAM L16

=> d scan

REGISTRY COPYRIGHT 2005 ACS on STN L17 2 ANSWERS

IN L-Phenylalanine, N-[(1,1-dimethylethoxy)carbonyl]-2-methoxy-N-methyl-5-[(phenylmethoxy)carbonyl]-, methyl ester (9CI)

MF C25 H31 N O7 Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L17 2 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN 1,2,4-Triazine-4(1H)-acetic acid, tetrahydro- α -(1-methylethyl)-3-oxo-1-[(phenylmethoxy)carbonyl]-, methyl ester, (α S)- (9CI)

MF C17 H23 N3 O5

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

=> file caplus COST IN U.S. DOLLARS SINCE FILE TOTAL **ENTRY** SESSION FULL ESTIMATED COST 0.86 214.68 SINCE FILE DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -2.92

FILE 'CAPLUS' ENTERED AT 12:15:49 ON 26 APR 2005
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FILE COVERS 1907 - 26 Apr 2005 VOL 142 ISS 18 FILE LAST UPDATED: 25 Apr 2005 (20050425/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 117 L18

4 L17

=> d 118 1-4 ti

- L18 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of peptide analogs as retroviral protease inhibitors
- L18 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of peptide analogs as retroviral protease inhibitors
- L18 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation and formulation of N-(α -aminoacyl)diaminohydroxyalkanes as HIV protease inhibitors
- L18 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Syntheses of peptide alkaloids, IX. Amino acids and peptides, XLVI. Total synthesis of mucronin B

=> d his

(FILE 'HOME' ENTERED AT 10:44:00 ON 26 APR 2005)

FILE 'CAPLUS' ENTERED AT 10:44:33 ON 26 APR 2005

L1 108 PHENYALANINE

L2 73592 PHENYLALANINE

L3 111423 DIABET?

L4 553 L2 AND L3

L5 0 LOGOFDF HOLD

FILE 'REGISTRY' ENTERED AT 10:53:30 ON 26 APR 2005

L6 STRUCTURE UPLOADED

L7 11 SEARCH L6 SSS SAM

FILE 'CAPLUS' ENTERED AT 10:54:54 ON 26 APR 2005

L8 18 L7

L9 0 L3 AND L8

FILE 'REGISTRY' ENTERED AT 10:55:45 ON 26 APR 2005 L10 228 SEARCH L6 SSS FULL

FILE 'CAPLUS' ENTERED AT 10:57:43 ON 26 APR 2005

L11 337 L10 L12 0 L3 AND L11 L13 875 PHENYL ACETIC L14 8713 PHENYLACETIC L15 107 L3 AND L14

FILE 'REGISTRY' ENTERED AT 12:14:55 ON 26 APR 2005

L16 STRUCTURE UPLOADED L17 2 SEARCH L16 SSS SAM

FILE 'CAPLUS' ENTERED AT 12:15:49 ON 26 APR 2005

L18 4 L17

=> 13 and 118

L19 0 L3 AND L18

=> file reg

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 2.18 216.86 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -2.92

FILE 'REGISTRY' ENTERED AT 12:16:44 ON 26 APR 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 American Chemical Society (ACS)

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TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

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Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=> search 116 sss full FULL SEARCH INITIATED 12:17:03 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - >1,000,000 TO ITERATE < 24.5% PROCESSED 320813 ITERATIONS

1171 ANSWERS

1363 ANSWERS

< 30.5% PROCESSED 400000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.29

FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**

BATCH **INCOMPLETE**

PROJECTED ITERATIONS:

EXCEEDS 1000000

PROJECTED ANSWERS:

EXCEEDS 4269

L20 1363 SEA SSS FUL L16

=> file caplus

COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST
161.76
378.62

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

SINCE FILE TOTAL
ENTRY SESSION
CA SUBSCRIBER PRICE

0.00 -2.92

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 120

L21 206 L20

=> 13 and 121

L22 35 L3 AND L21

=> d 122 25-35 ti

- L22 ANSWER 25 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of 3-(oxazolylalkoxyphenyl)propionic acids and analogs as modulators of peroxisome proliferator activated receptors for treatment of diabetes and related conditions
- L22 ANSWER 26 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of benzyloxyphenyloxobutyrates and related compounds for the treatment of metabolic disorders

- L22 ANSWER 27 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of pyrrolidine, piperidine, or piperazine amino acid derivatives as melanocortin receptor ligands
- L22 ANSWER 28 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Dipeptidyl peptidase inhibitors for the treatment or prevention of diabetes
- L22 ANSWER 29 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of indolizinylpyrrole derivates for treating cytokine mediated diseases
- L22 ANSWER 30 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of phenylalkanoic acid derivatives as peroxisome proliferator activated receptor (PPAR) agonists
- L22 ANSWER 31 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of phenylalkenoic acid derivatives and phenylalkanoic acid derivatives as PPAR α agonists for treatment of hyperlipidemia, arteriosclerosis, obesity, and **diabetes**
- L22 ANSWER 32 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of (phenymethyl)alkanoic acid derivatives as PPARα agonists for treatment of arteriosclerosis, obesity, diabetes, etc.
- L22 ANSWER 33 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of (phenylmethyl)alkanoic acid derivatives as PPAR α (peroxisome proliferator-activated receptor α) agonists useful in treatment of hyperlipidemia, arteriosclerosis, obesity, and diabetes
- L22 ANSWER 34 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of phenylmethylalkanoic acid derivatives as PPAR α agonists useful in the treatment of hyperlipidemia, arteriosclerosis, diabetes, and obesity
- L22 ANSWER 35 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Enantio-dependent binding and transactivation of optically active phenylpropanoic acid derivatives at human peroxisome proliferator-activated receptor alpha
- => d 122 25,26,31,32-35 ti fbib abs
- L22 ANSWER 25 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of 3-(oxazolylalkoxyphenyl)propionic acids and analogs as modulators of peroxisome proliferator activated receptors for treatment of diabetes and related conditions
- AN 2002:964190 CAPLUS
- DN 138:39272
- TI Preparation of 3-(oxazolylalkoxyphenyl)propionic acids and analogs as modulators of peroxisome proliferator activated receptors for treatment of diabetes and related conditions
- IN Gossett, Lynn Stacy; Green, Jonathan Edward; Henry, James Robert; Jones, Winton Dennis, Jr.; Matthews, Donald Paul; Shen, Quan Rong; Smith, Daryl Lynn; Vance, Jennifer Ann; Warshawsky, Alan M.
- PA Eli Lilly and Company, USA
- SO PCT Int. Appl., 438 pp. CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 1

	PATENT NO.				KIND DATE			APPLICATION NO.					DATE					
ΡI	WO	2002	1004	03		A1 20021219			WO 2002-US15143						2	0020	 524	
		W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,
			CN,	co,	CR,	CU,	CZ,	CZ,	DE,	DK,	DK,	DM,	DZ,	EC,	EE,	EE,	ES,	FI,
			FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,
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			MZ,	NO,	ΝŻ,	OM,	PH,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SK,	SL,
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				BY,														
		RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,
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	ΝZ	5295	50			Α		2003	1219								0020	
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	EΡ	1401	434			A 1		2004	0331		EP 2	2002-	7463	80		2	0020	524
		R:	-		•			-		•		IT,	LI,	LU,	NL,	SE,	MC,	PT,
			ΙE,	SI,	LT,	LV,	FΙ,	RO,	MK,									
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	BR	2002	0101	67		Α		2004	0406			2002-				_	0020	
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						2001-					0010							
									WO 2	2002-	0212	143	1	w 2	0020	524		
OS MARPAT 138:39272																		

GI

AB Title compds. I [wherein n = 2-5; V = a bond or O; X = CH2 or O; p = 0 or 1; m = 1-4; Y1 = (un)substituted (hetero)aryl; Y2 and Y3 = independently H, alkyl, or alkoxy; Y4 = (un)substituted alk(en/yn)ylaminoalkyl, carboxyaminoalkyl, (thio)ureidoalkyl, carbamoylalkyl, aminoalkyl,

alkoxyalkyl, alkylthioalkyl, or CN; R5 = H or alkyl; and pharmaceutically acceptable salts, solvates, hydrates, or stereoisomers thereof] were prepared as peroxisome proliferator activated receptor (PPAR) modulators (no data). For example, 3-[2-(1,3-dioxo-1,3-dihydroisoindolo-2-ylmethyl)-4-hydroxyphenyl]propionic acid tert-Bu ester was coupled with toluene-4-sulfonic acid 2-(5-methyl-2-phenyloxazol-4-yl)ethyl ester in the presence of Cs2CO3 in DMF. Deprotection of the amine using NaBH4 in isopropanol followed by conversion to the carbamate and deesterification gave II. I are useful for the treatment of Syndrome X, Type II diabetes, hyperglycemia, hyperlipidemia, obesity, coagulopathy, hypertension, arteriosclerosis, and other disorders related to Syndrome X, as well as cardiovascular diseases (no data).

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L22 ANSWER 26 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of benzyloxyphenyloxobutyrates and related compounds for the treatment of metabolic disorders
- AN 2002:964135 CAPLUS
- DN 138:24543
- TI Preparation of benzyloxyphenyloxobutyrates and related compounds for the treatment of metabolic disorders
- IN Sharma, Shalini; Von Borstel, Reid W.; Hodge, Kirvin L.
- PA Wellstat Therapeutics Corporation, USA; Bamat, Michael K.
- SO PCT Int. Appl., 242 pp. CODEN: PIXXD2
- DT Patent
- LA English
- EAN CMT 1

FAN.						KIND DATE			APPLICATION NO.						DATE				
PI	_	2002				A2 A3		2002 2004		1	WO 2	2002-1	JS18	388		2	20020	612	
		₩:					AT,	AU,	ΑZ,			BG, EE,							
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			-	-	-	-						, MW,		-					•
			FL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK	, SL,	ТJ,	TM,	TN,	18,	TT,	TZ,	•
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		RW:										, TZ,							
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			GN,	GQ,	GW,	ML,	MK,	NE,	SN,			2001-:	2072	0 û D		n ,	20010	612	
	115	2003	1/01	07		λ1		2003	0807								20010		
	US 2003149107 EP 1461323			ΑI		2003	0007			2002-					20020				
				A2		2004	0929			2002-					20020				
		R:	AT,	BE, FI,	CH,	DE,						, IT,			NL,				
			•	•	•						US 2	2001-	2972	82P		P 2	20010	612	
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	JР	2005	5010	12		Т2		2005	0113			2003-					20020		
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	US	2004	0778	96		A1		2004	0422			2003-					20031		
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	110	2004	0025	10		A1		2004	0512			2002-: 2003-			4		20020		
	0.5	2004	0923	10		ΑI		2004	0313			2003-1 2001-1					20031		
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	US	2004	0925	16		A1		2004	0513			2003-			•		20031		
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				US	2001-297282P	P	20010612
	•			US	2002-167839	A3	20020612
US	2004236100	A1	20041125	US	2003-684660		20031014
US	6858602	B2	20050222				
				US	2001-297282P	P	20010612
				US	2002-167839	A3	20020612
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				US	2001-297282P	P	20010612
				US	2002-167839	A3	20020612
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				US	2001-297282P	P	20010612
				US	2002-167839	A1	20020612
US	2005004115	A1	20050106	US	2004-892950		20040716
				US	2001-297282P	P	20010612
				US	2002-167839	А3	20020612
				US	2003-685183	A3	20031014
3 / 3 T	NAM 100.04540						

OS MARPAT 138:24543 GΙ

$$A(CH_2)_p(NR^5)_q(CH_2)_nO$$
(CH₂)_mCOXCOQ I

Biol. active title compds. [I; n = 1, 2; m, q, p = 0, 1; R5 = alkyl; R9 = alkylAB H, halo, alkoxy; A = (halo-, alkyl-, perfluoromethyl-, alkoxy-, perfluoromethoxy-substituted) Ph, (Me-, Et-substituted) cycloalkyl, 5-6 membered heteroarom. ring having 1-2 N, S, O atoms; X = CH2, Q = OR1, R1 =Et; or X = CH2CR12R13, CH2CH(NHAc), Q = OR1, R1 = H, alkyl; or X = CH2CH2, Q = NR10R11; R12, R13 = H, Me; 1 of R10, R11 = H, alkyl, OH, the other = H, alkyl], were prepared Thus, 4-(2-fluorobenzyloxy) acetophenone (preparation given) in THF and DMPU was treated with a solution of Li bis(trimethylsilyl)amide at -60°; after 10 min, tert-Bu bromoacetate was added followed by stirring for an addnl. 10 min and warming to room temperature for 4 h to give tert-Bu 4-[4-(2fluorobenzyloxy)phenyl]-4-oxobutyrate. The latter was stirred with CF3CO2H in CH2Cl2 to give 4-[4-(2-fluorobenzyloxy)phenyl]-4-oxobutyric acid. Tested I showed antidiabetic activity in a variety of tests. I are useful in treatment of various metabolic disorders such as insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis and arteriosclerosis.

L22 ANSWER 31 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN

TΙ Preparation of phenylalkenoic acid derivatives and phenylalkanoic acid derivatives as PPARa agonists for treatment of hyperlipidemia, arteriosclerosis, obesity, and diabetes

2002:428860 CAPLUS AN

DN 137:5999

Preparation of phenylalkenoic acid derivatives and phenylalkanoic acid ΤI derivatives as PPARa agonists for treatment of hyperlipidemia, arteriosclerosis, obesity, and diabetes Miyachi, Hiroyuki; Tanase, Takahiro; Murakami, Kouji

IN

Kyorin Pharmaceutical Co., Ltd., Japan PA

SO PCT Int. Appl., 53 pp. CODEN: PIXXD2

DTPatent

LA Japanese

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

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WO 2002044131
                               A1
                                      20020606
                                                     WO 2001-JP10354
PΙ
                                                                                 20011128
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          W:
               CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
               GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
               LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
               PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
               BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                                     JP 2000-363678
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                               Α5
                                      20020611
     AU 2002022551
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                                                                                 20011128
                                                     JP 2000-363678
                                                                             A 20001129
                                                     WO 2001-JP10354
                                                                             W 20011128
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OS MARPAT 137:5999

GΙ

$$R^2$$
 N
 MeO
 $(CH_2)_n-C=C$
 R^2

AB The title compds. I [n is 0 or 1; R1 represents hydrogen or lower alkyl; R2 represents carboxyl, lower alkoxycarbonyl, carbamoyl, hydroxyaminocarbonyl, lower alkoxyaminocarbonyl or 5-tetrazolyl; and the dotted line shows together with the solid line a double bond or a single bond; a proviso is given] are prepared For example, 3-[3-[N-[[4-(trifluoromethyl)phenyl]methyl]carbamoyl]-4-methoxyphenyl]-2-ethyl-2propenoic acid (II) was prepared The effect of II on peroxisome proliferator-activated receptors α (PPAR α) was demonstrated.

Ι

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 32 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN L22

Preparation of (phenymethyl)alkanoic acid derivatives as PPARa ΤI agonists for treatment of arteriosclerosis, obesity, diabetes, etc.

AN 2002:428859 CAPLUS

DN 137:5998

ΤI Preparation of (phenymethyl)alkanoic acid derivatives as PPARa agonists for treatment of arteriosclerosis, obesity, diabetes,

Miyachi, Hiroyuki; Nomura, Masahiro; Takahashi, Yukie; Tanase, Takahiro; IN Murakami, Kouji

Kyorin Pharmaceutical Co., Ltd., Japan PΑ

SO PCT Int. Appl., 52 pp.

CODEN: PIXXD2

DΤ Patent

LΑ Japanese

FAN.CNT 1

	PATENT NO.				KIND DATE				APPLICATION NO.					DATE				
PI	WO 2002044130			A1 20020606			WO 2001-JP10353						20011128					
		W:	ΑE,	AG,	ΑL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
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			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
			PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	ΤZ,	UA,

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UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
    RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
        CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
        BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                       JP 2000-363677
                                                           A 20001129
                     A5
                           20020611
                                       AU 2002-22550
AU 2002022550
                                                               20011128
                                       JP 2000-363677
                                                              20001129
                                       WO 2001-JP10353
                                                           W
                                                              20011128
MARPAT 137:5998
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AB The title compds. I [R1 represents hydrogen, halogeno, hydroxy, 2-phenylethyl, 2-phenylethoxy, hydroxyphenoxy or benzyloxyphenoxy; and R2 represents lower (C1-4) alkyl] are prepared I are lipid-lowering drugs (particularly in the liver), drugs preventing the progress of arteriosclerosis, anti-obesity drugs and remedies for diabetes. For example, 2-[[3-[N-[(4-chlorophenyl)methyl]carbamoyl]-4-methoxyphenyl]methyl]butyric acid (II) was prepared The PPARα agonist activity of II was demonstrated.

Ι

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L22 ANSWER 33 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN

TI Preparation of (phenylmethyl)alkanoic acid derivatives as PPAR α (peroxisome proliferator-activated receptor α) agonists useful in treatment of hyperlipidemia, arteriosclerosis, obesity, and diabetes

AN 2002:428858 CAPLUS

DN 137:5997

OS GI

TI Preparation of (phenylmethyl)alkanoic acid derivatives as PPAR α (peroxisome proliferator-activated receptor α) agonists useful in treatment of hyperlipidemia, arteriosclerosis, obesity, and

diabetes

IN Miyachi, Hiroyuki; Murakami, Kouji

PA Kyorin Pharmaceutical Co., Ltd., Japan

SO PCT Int. Appl., 33 pp.

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

PATENT 1	NO.			KIN	D	DATE		. 1	APPL.	ICAT:	I NOI	. O <i>i</i>		D?	\ΤΕ	
WO 2002044129			A1 20020606			WO 2001-JP10352						20011128				
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
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RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑT,	BE,	CH,
	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,
	WO 2002 W:	W: AE, CO, GM, LS, PL, UG, RW: GH,	WO 2002044129 W: AE, AG, CO, CR, GM, HR, LS, LT, PL, PT, UG, US, RW: GH, GM,	WO 2002044129 W: AE, AG, AL, CO, CR, CU, GM, HR, HU, LS, LT, LU, PL, PT, RO, UG, US, UZ, RW: GH, GM, KE,	WO 2002044129 A1 W: AE, AG, AL, AM, CO, CR, CU, CZ, GM, HR, HU, ID, LS, LT, LU, LV, PL, PT, RO, RU, UG, US, UZ, VN, RW: GH, GM, KE, LS,	WO 2002044129 A1 W: AE, AG, AL, AM, AT, CO, CR, CU, CZ, DE, GM, HR, HU, ID, IL, LS, LT, LU, LV, MA, PL, PT, RO, RU, SD, UG, US, UZ, VN, YU, RW: GH, GM, KE, LS, MW,	WO 2002044129 A1 20020 W: AE, AG, AL, AM, AT, AU, CO, CR, CU, CZ, DE, DK, GM, HR, HU, ID, IL, IN, LS, LT, LU, LV, MA, MD, PL, PT, RO, RU, SD, SE, UG, US, UZ, VN, YU, ZA, RW: GH, GM, KE, LS, MW, MZ,	WO 2002044129 A1 20020606 W: AE, AG, AL, AM, AT, AU, AZ, CO, CR, CU, CZ, DE, DK, DM, GM, HR, HU, ID, IL, IN, IS, LS, LT, LU, LV, MA, MD, MG, PL, PT, RO, RU, SD, SE, SG, UG, US, UZ, VN, YU, ZA, ZW, RW: GH, GM, KE, LS, MW, MZ, SD,	WO 2002044129 W: AE, AG, AL, AM, AT, AU, AZ, BA, CO, CR, CU, CZ, DE, DK, DM, DZ, GM, HR, HU, ID, IL, IN, IS, JP, LS, LT, LU, LV, MA, MD, MG, MK, PL, PT, RO, RU, SD, SE, SG, SI, UG, US, UZ, VN, YU, ZA, ZW, AM, RW: GH, GM, KE, LS, MW, MZ, SD, SL,	WO 2002044129 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, GM, HR, HU, ID, IL, IN, IS, JP, KE, LS, LT, LU, LV, MA, MD, MG, MK, MN, PL, PT, RO, RU, SD, SE, SG, SI, SK, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ,	WO 2002044129 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ,	WO 2002044129 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG,	WO 2002044129 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM,	WO 2002044129 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW,	WO 2002044129 A1 20020606 WO 2001-JP10352 20 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT,	WO 2002044129 A1 20020606 WO 2001-JP10352 200113

BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

JP 2000-363676 A 20001129

AU 2002022549 A5 20020611 AU 2002-22549 20011128

JP 2000-363676 A 20001129

WO 2001-JP10352 W 20011128

OS MARPAT 137:5997

GI

$$F_3C$$
 (CH₂) $n-N$ R Et

AB The title compds. I [n is 0, 1 or 2; and R represents hydrogen or lower (C1-10) alkyl in case where n is 0 or 2, or lower (C1-10) alkyl in case where n is 1] are prepared For example, I [n = 2; R = H] (II) was prepared The effect of II on the peroxisome proliferator-activated receptor α was demonstrated.

Ι

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L22 ANSWER 34 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN

TI Preparation of phenylmethylalkanoic acid derivatives as PPARα agonists useful in the treatment of hyperlipidemia, arteriosclerosis, diabetes, and obesity

AN 2002:428856 CAPLUS

DN 137:20225

Preparation of phenylmethylalkanoic acid derivatives as PPAR α agonists useful in the treatment of hyperlipidemia, arteriosclerosis, diabetes, and obesity

IN Mıyachi, Hiroyuki; Nomura, Masahiro; Murakami, Kouji

PA Kyorin Pharmaceutical Co., Ltd., Japan

SO PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

	PATENT NO.				KIND DATE				APPLICATION NO.						DATE			
PI	WO 2002044127			A1 20020606			WO 2001-JP10355						20011128					
	W:	ΑE,	AG,	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	
		UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM	
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,	
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG	
									•	JP 2	000-	3636	79	2	A 2	0001	129	
	AU 2002	0225	52		A 5		2002	0611		AU 2	002-	2255	2		2	0011	128	
									1	JP 2	000-	3636	79	2	A 2	0001	129	
									1	WO 2	001-	JP10	355	1	₩ 2	0011	128	

OS MARPAT 137:20225

GI

$$R^{1}$$
 $(CH_{2})_{n}-A$
 $X-C-CO-OR^{5}$
 R^{3}

AB The title compds. I [R1 represents trifluoromethyl, optionally substituted phenoxy, etc.; R2 represents hydrogen or lower alkoxy; R3, R4 and R5 represent each hydrogen or lower alkyl; A represents NHCO or CONH; X is located at the para-position relative to A and represents oxygen or sulfur, or X is located at the para-position relative to R2 and represents oxygen or sulfur; and n is an integer of from 0 to 2], useful as PPARα agonists (no data) for the treatment of hyperlipidemia, arteriosclerosis, diabetes, and obesity, are prepared For example, 2-[[4-[N-[[4-(trifluoromethyl)phenyl]methyl]carbamoyl]-3-methoxyphenyl]methyl]butyric acid was prepared

Ι

- RE.CNT 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L22 ANSWER 35 OF 35 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Enantio-dependent binding and transactivation of optically active phenylpropanoic acid derivatives at human peroxisome proliferator-activated receptor alpha
- AN 2002:97666 CAPLUS
- DN 137:78739
- TI Enantio-dependent binding and transactivation of optically active phenylpropanoic acid derivatives at human peroxisome proliferator-activated receptor alpha
- AU Miyachi, Hiroyuki; Nomura, Masahiro; Tanase, Takahiro; Suzuki, Masahiro; Murakami, Koji; Awano, Katsuya
- CS Kyorin Pharmaceutical Co., Ltd., Discovery Research Laboratories, Shimotsuga-gun, Nogi-machi, Tochigi, 329-0114, Japan
- SO Bioorganic & Medicinal Chemistry Letters (2002), 12(3), 333-335 CODEN: BMCLE8; ISSN: 0960-894X
- PB Elsevier Science Ltd.
- DT Journal
- LA English
- OS CASREACT 137:78739

GΙ

$$_{\mathrm{F3C}}$$
 $_{\mathrm{Me}}^{\mathrm{CO}_{2}\mathrm{H}}$

AB Optically active phenylpropanoic acid derivs. [(S)-I, and (R)-I] were prepared, and their affinities for peroxisome proliferator-activated receptor (PPAR) α and PPAR γ were evaluated. Binding assay and cell-based reporter assay indicated that the activity of these compds. is enantio-dependent, and resides exclusively on the (S)-isomer.

Ι

RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> logoff hold COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY-SESSION FULL ESTIMATED COST 26.12 404.74 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -5.11-8.03

SESSION WILL BE HELD FOR 60 MINUTES STN INTERNATIONAL SESSION SUSPENDED AT 12:23:14 ON 26 APR 2005

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID:SSSPTA1623PAZ

PASSWORD:

* * * * * * RECONNECTED TO STN INTERNATIONAL * * * * * SESSION RESUMED IN FILE 'CAPLUS' AT 12:24:46 ON 26 APR 2005 FILE 'CAPLUS' ENTERED AT 12:24:46 ON 26 APR 2005 COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S. DOLLARS FULL ESTIMATED COST	SINCE FILE ENTRY 26.12	TOTAL SESSION 404.74
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) CA SUBSCRIBER PRICE	SINCE FILE ENTRY -5.11	TOTAL SESSION -8.03
=> file reg COST IN U.S. DOLLARS FULL ESTIMATED COST	SINCE FILE ENTRY 26.12	TOTAL SESSION 404.74
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) CA SUBSCRIBER PRICE	SINCE FILE ENTRY -5.11	TOTAL SESSION -8.03

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STRUCTURE FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0 DICTIONARY FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

=>

Uploading C:\Documents and Settings\PZucker\My Documents\Examination Auxillary files\10797458\10797458 modified subgenus 3.str

chain nodes :

1 2 3 4 5 6 7 8 10 11 12 17 18

chain bonds :

1-2 1-8 2-3 3-4 4-5 4-10 5-6 6-7 7-11 11-12 12-17 12-18

exact/norm bonds :

2-3 3-4 4-10 5-6 6-7 12-17 12-18

exact bonds :

1-2 1-8 4-5 7-11 11-12

G1:C,O,S,N,SO2

Match level:

1:Atom 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:Atom 7:CLASS 8:Atom 10:CLASS

11:CLASS 12:CLASS 17:CLASS 18:CLASS

Generic attributes :

6:

Number of Carbon Atoms : less than 7 Type of Ring System : Monocyclic

L23 STRUCTURE UPLOADED

=> d 123 L23 HAS NO ANSWERS L23 STR

G1 C,O,S,N,SO2

Structure attributes must be viewed using STN Express query preparation.

6 ANSWERS

=> search 123 sss sam
SAMPLE SEARCH INITIATED 12:25:18 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 136104 TO ITERATE

0.7% PROCESSED 1000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**

BATCH **INCOMPLETE**

PROJECTED ITERATIONS: EXCEEDS 1000000

PROJECTED ANSWERS: EXCEEDS 14618

L24 6 SEA SSS SAM L23

=> d scan

L24 6 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN L-Threonine, L-α-aspartyl-L-prolyl-L-valyl-L-threonyl-L-leucyl-L-asparaginyl-L-valyl- (9CI)

SQL 8

MF C37 H63 N9 O14

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):6

L24 6 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN Benzeneacetic acid, 2-(1-oxoundecyl)- (9CI)

MF C19 H28 O3

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L24 6 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN L-Serine, glycyl-L-valyl-L-leucyl-L-prolyl-L-alanyl-L-leucyl-L-prolyl-L-glutaminyl-L-valyl-L-cysteinyl-L-leucyl-L-threonyl-L-cysteinyl-L-α-aspartyl-L-prolyl-L-arginyl-L-phenylalanyl-L-glutaminyl-L-α-aspartyl-L-seryl- (9CI)

SQL 23

MF C106 H170 N28 O35 S2

PAGE 1-C

PAGE 2-C

L24

6 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN L-Phenylalanine, 4,4'-[2,6-pyridinediylbis[imino[(1S)-1-(3-methoxy-3-oxopropyl)-2-oxo-2,1-ethanediyl]iminocarbonyl]]bis[N-[(1,1-IN dimethylethoxy)carbonyl]- (9CI)

C47 H59 N7 O16 MF

PAGE 1-B

L24 6 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

IN L-Threonine, L-prolyl-L-alanyl-L-seryl-L-arginyl-L-isoleucyl-L- α -aspartyl-L-isoleucyl-L- α -aspartyl-L-prolyl- (9CI)

SQL 10

MF C46 H77 N13 O17

L24 6 ANSWERS REGISTRY COPYRIGHT 2005 ACS on STN

Labeled Levaly1-L-proly1-L- α -aspar phenylalany1-L- α -glutamy1-L-sery1- (9CI) 9 $L\text{--Isoleucine, L--valyl-L--prolyl-L--a-aspartyl-L--prolyl-L--valyl-L--prol$ IN

SQL

MF C47 H71 N9 O15

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

=> search 123 sss full FULL SEARCH INITIATED 12:25:54 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - >1,000,000 TO ITERATE

< 7.5% PROCESSED 203829 ITERATIONS 1574 ANSWERS

< 11.9% PROCESSED 324370 ITERATIONS 2498 ANSWERS

< 14.7% PROCESSED 400000 ITERATIONS 3057 ANSWERS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.52

FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**

BATCH **INCOMPLETE**

PROJECTED ITERATIONS: EXCEEDS 1000000

PROJECTED ANSWERS: EXCEEDS 20346

L25 3057 SEA SSS FUL L23

=> file caplus

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST 162.19 566.93

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL

ENTRY SESSION

CA SUBSCRIBER PRICE 0.00 -8.03

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FILE COVERS 1907 - 26 Apr 2005 VOL 142 ISS 18 FILE LAST UPDATED: 25 Apr 2005 (20050425/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> 125

L26 762 L25

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=> 13 and 126
L27 57 L3 AND L26
=> d his
     (FILE 'HOME' ENTERED AT 10:44:00 ON 26 APR 2005)
    FILE 'CAPLUS' ENTERED AT 10:44:33 ON 26 APR 2005
L1
          108 PHENYALANINE
L2
         73592 PHENYLALANINE
        111423 DIABET?
L3
           553 L2 AND L3
L4
L5
             0 LOGOFDF HOLD
   FILE 'REGISTRY' ENTERED AT 10:53:30 ON 26 APR 2005
               STRUCTURE UPLOADED
L6
            11 SEARCH L6 SSS SAM
L7
    FILE 'CAPLUS' ENTERED AT 10:54:54 ON 26 APR 2005
            18 L7
r_8
             0 L3 AND L8
L9
     FILE 'REGISTRY' ENTERED AT 10:55:45 ON 26 APR 2005
L10
           228 SEARCH L6 SSS FULL
    FILE 'CAPLUS' ENTERED AT 10:57:43 ON 26 APR 2005
          337 L10
L11
             0 L3 AND L11
L12
L13
           875 PHENYL ACETIC
L14
          8713 PHENYLACETIC
           107 L3 AND L14
L15
    FILE 'REGISTRY' ENTERED AT 12:14:55 ON 26 APR 2005
L16
             STRUCTURE UPLOADED
L17
             2 SEARCH L16 SSS SAM
 .. FILE 'CAPLUS' ENTERED AT 12:15:49 ON 26 APR 2005
     4 L17
L18
             0 L3 AND L18
L19
     FILE 'REGISTRY' ENTERED AT 12:16:44 ON 26 APR 2005
L20
          1363 SEARCH L16 SSS FULL
    FILE 'CAPLUS' ENTERED AT 12:17:40 ON 26 APR 2005
L21
          206 L20
L22
            35 L3 AND L21
   FILE 'REGISTRY' ENTERED AT 12:24:53 ON 26 APR 2005
L23
              STRUCTURE UPLOADED
L24
             6 SEARCH L23 SSS SAM
L25
        3057 SEARCH L23 SSS FULL
     FILE 'CAPLUS' ENTERED AT 12:26:54 ON 26 APR 2005
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=> 127 not 122 L28 55 L27 NOT L22

762 L25

57 L3 AND L26

=> d 128 45-55 ti

L26

L27

L28 ANSWER 45 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
TI Preparation of phenoxyalkanamides as amide linker peroxisome proliferator

activated receptor agonists for treating and/or preventing diabetes mellitus and syndrome X

- L28 ANSWER 46 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Insulin and IGF-1 receptor peptide agonists and antagonists, and therapeutic use
- L28 ANSWER 47 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Expression of Preproinsulin-2 Gene Shapes the Immune Response to Preproinsulin in Normal Mice
- L28 ANSWER 48 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Pancreatitis—associated protein and methods for promoting $\beta\text{-cell}$ neogenesis
- L28 ANSWER 49 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Targets for therapeutic intervention identified in the human mitochondrial proteome
- L28 ANSWER 50 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Active antiangiogenic therapy
- L28 ANSWER 51 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation and compositions of nitrosothio (hetero)cyclic nitric oxide donors
- L28 ANSWER 52 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Insulin and IGF-1 receptor peptide agonists and antagonists, and therapeutic use
- L28 ANSWER 53 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Chemokine binding peptides capable of modulating the biological activity of chemokines, and therapeutic use
- L28 ANSWER 54 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Hsp70-derived peptides an uses thereof in the diagnosis and treatment of autoimmune diseases
- L28 ANSWER 55 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Non-invasive measurement of metabolism of biological molecules not easily accessible to direct sampling by label incorporation into metabolites and catabolites

=> d 128 45 ti fbib abs

- L28 ANSWER 45 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Preparation of phenoxyalkanamides as amide linker peroxisome proliferator activated receptor agonists for treating and/or preventing diabetes mellitus and syndrome X
- AN 2004:2837 CAPLUS
- DN 140:59411
- TI Preparation of phenoxyalkanamides as amide linker peroxisome proliferator activated receptor agonists for treating and/or preventing diabetes mellitus and syndrome X
- IN Ferritto Crespo, Rafael; Martin, Jose Alfredo; Martin-Ortega, Finger Maria
 Dolores; Rojo Garcia, Isabel; Shen, Quanrong; Warshawsky, Alan M.; Xu,
 Yanping
- PA Eli Lilly and Company, USA
- SO PCT Int. Appl., 168 pp. CODEN: PIXXD2
- DT Patent
- LA English

MARPAT 140:59411

GΙ

AB The present invention is directed to phenoxyalkanamides (shown as I; variables defined below; e.g. II), compns., and their use as peroxisome proliferator activated receptor agonists for treating and/or preventing diabetes mellitus and syndrome X. The binding and cotransfection efficacy values found for compds. of this invention that are useful for modulating a PPAR α receptor are about <100 nM and >50%, resp. Although the methods of preparation are not claimed, .apprx.140 example prepns. of I are included. For example, II was prepared in 3 steps starting from (2S)-2-ethoxy-3-(4-hydroxyphenyl)propionic acid Me ester, (2S)-2-hydroxypropionic acid benzyl ester and involving intermediates (2S)-3-[4-[(1R)-1-[(benzyloxy)carbonyl]ethyl]oxy]phenyl]-2ethoxypropionic acid Et ester and (2S)-3-[4-[((1R)-1carboxyethyl)oxy]phenyl]-2-ethoxypropionic acid. For I: R1 = H, C1-C8 alkyl, C3-C6 cycloalkyl, aryl-C0-4-alkyl, heteroaryl-C0-4-alkyl, aminoC1-C4alkyl, C3-C6 cycloalkylaryl-C0-2-alkyl, arylheteroC1-C8alkyl, -CHC(O)C1-C4 alkoxy, C0-4-alkyl-C(O)heteroC1-C8alkyl, and -CH2C(O)-R15R16. R2 = C1-C8 alkyl, C3-C6 cycloalkyl, aryl-C0-C4-alkyl, heteroaryl-C0-C4alkyl, heteroC1-C6cycloalkylaryl, heteroC1-C6cycloalkylarylC1-C4alkyl, aminoC1-C4alkyl, C3-C6 cycloalkylaryl-C0-C2-alkyl, arylheteroC1-C8alkyl, C0-C4-alkyl-C(0)heteroC1-C8alkyl, -CH(C(0)OCH3)benzyl, and

-CH2C(O)R15''R16''. R1 and R2 together may form a heterocyclic ring which heterocyclic ring is (un)substituted with 1-3 substituents R1' and which heterocyclic ring is optionally fused with an aryl; E = C(R3)(R4)A, (CH2)nCOOR13, aryl-C0-C4-alkyl, thio-C1-C4-alkyl, thioaryl, arylC1-C4alkoxy, C1-C4alkoxy C1-C4alkyl, aminoaryl, and aminoC1-C4alkyl. R5 and R6 = H, C1-C8 alkyl, aryl-C0-C4-alkyl, heteroaryl-C0-C4-alkyl, C3-C6 cycloalkyl, aryl-C0-C2-alkyl, C3-C6 cycloalkyl, aryl-C0-C2-alkyl, C3-C6 cycloalkyl-C0-2-alkyl, and -CH2C(O)R17R18.

RE.CNT 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d 128 34-44 ti

- L28 ANSWER 34 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Protein tyrosine phosphatase inhibitors
- L28 ANSWER 35 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Nogo receptor antagonists for promoting survival of neuron and treating multiple sclerosis, CNS neuropathy, and traumatic brain or spinal cord injury
- L28 ANSWER 36 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Methods for the production and therapeutic uses of cytokine receptor INSP076 and ligands
- L28 ANSWER 37 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Cell penetrating peptides
- L28 ANSWER 38 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Cell penetrating peptides
- L28 ANSWER 39 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Cell penetrating peptides
- L28 ANSWER 40 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Cell penetrating peptides
- L28 ANSWER 41 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Cell penetrating peptides
- L28 ANSWER 42 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Cell penetrating peptides
- L28 ANSWER 43 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Use of peptides derived from junctional adhesion molecules to permeabilize mucosa for improved efficiency of mucosal delivery of therapeutic compounds
- L28 ANSWER 44 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Epitopes of viral LMP1 and LMP2 proteins for inducing tolerance to target antigens and for treating allergy, autoimmune disease and transplant rejection

=> d 128 34-36 ti fbib abs

- L28 ANSWER 34 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
- TI Protein tyrosine phosphatase inhibitors
- AN 2004:203856 CAPLUS
- DN 140:247109
- TI Protein tyrosine phosphatase inhibitors
- IN Hooft van Huijsduijnen, Rob; Walchli, Sebastien; Arigoni, Fabrizio

```
Applied Research Systems Ars Holding N.V., Neth. Antilles
PΑ
     PCT Int. Appl., 76 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
     PATENT NO.
                           KIND
                                   DATE
                                                APPLICATION NO.
                                                                         DATE
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     _____
     WO 2004020466
                                   20040311
                                                WO 2003-EP50385
                                                                         20030820
PΙ
                            A1
                                   20050120
     WO 2004020466
                           C1
              AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
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                                                EP 2002-19357
                                                                    A 20020829
os
     MARPAT 140:247109
AB
     The invention relates to phosphopeptides inhibiting protein tyrosine
     phosphatases, and their medical uses. The invention relates to
     phosphopeptides and phosphopeptide derivs. inhibiting protein tyrosine
     phosphatases.
RE.CNT 4
               THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
               ALL CITATIONS AVAILABLE IN THE RE FORMAT
L28
     ANSWER 35 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
TI
     Nogo receptor antagonists for promoting survival of neuron and treating
     multiple sclerosis, CNS neuropathy, and traumatic brain or spinal cord
     injury
     2004:142908 CAPLUS
ΑN
     140:198086
DN
ΤI
     Nogo receptor antagonists for promoting survival of neuron and treating
     multiple sclerosis, CNS neuropathy, and traumatic brain or spinal cord
     injury
     Lee, Daniel H. S.; Pepinsky, R. Blake; Li, Weiwei; Rabacchi, Sylvia A.;
IN
     Relton, Jane K.; Worley, Dane S.; Strittmatter, Stephen M.; Sah, Dinah Y.
PA
     Yale University, USA; Biogen, Inc.
SO
     PCT Int. Appl., 133 pp.
     CODEN: PIXXD2
DT
     Patent
T.A
     English
FAN.CNT 1
     PATENT NO.
                           KIND
                                   DATE
                                                APPLICATION NO.
                                                                         DATE
PΙ
     WO 2004014311
                            A2
                                   20040219
                                                WO 2003-US25004
                                                                          20030807
                            A3
                                   20040429
     WO 2004014311
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
              PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
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              BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
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WO 2005016955

A2

20050224

US 2002-402866P P 20020810

20040130

WO 2004-US2702

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CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
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         NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE,
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             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                              WO 2003-US25004
                                                                  A · 20030807
     Disclosed are immunogenic Nogo receptor-1 polypeptides, Nogo receptor-1
     antibodies, antigen-binding fragments thereof, soluble Nogo receptors and
     fusion proteins thereof and nucleic acids encoding the same. Also
     disclosed are compns. comprising, and methods for making and using, such
     Nogo receptor antibodies, antigen-binding fragments, humanized and
     chimeric antibodies thereof, soluble Nogo receptors and fusion proteins
     thereof and nucleic acids or viral vector encoding the same for gene
     therapy. These Nogo receptor-1, antagonists are useful for inhibiting
     growth cone collapse of neuron, decreasing inhibition of neurite
     outgrowth, promoting survival of CNS neuron and axonal growth, and are
     therefore useful for treating multiple sclerosis, ALS, Huntington's
     disease, Alzheimer's disease, Parkinson's disease, diabetes
     neuropathy, stroke, traumatic brain injury or spinal cord injury.
     ANSWER 36 OF 55 CAPLUS COPYRIGHT 2005 ACS on STN
     Methods for the production and therapeutic uses of cytokine receptor
     INSP076 and ligands
     2004:60555 CAPLUS
     140:127207
     Methods for the production and therapeutic uses of cytokine receptor
     INSP076 and ligands
     Rodrigues, Tania Maria; Fagan, Richard Joseph; Phelps, Christopher
     Benjamin; Power, Christine
     Ares Trading S.A., Switz.
     PCT Int. Appl., 86 pp.
     CODEN: PIXXD2
     Patent
     English
FAN.CNT 1
                                              APPLICATION NO.
                          KIND
                                 DATE
     PATENT NO.
                                                                       DATE
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                                              _____
                                 20040122
                                              WO 2003-GB3107
     WO 2004007552
                          A1
                                                                       20030717
     WO 2004007552
                          C1
                                 20040415
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             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
             PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
             TR, TT, TZ, UA, UG, US, UZ; VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                                                   A 20020717
                                              GB 2002-16661
     The invention is based on the discovery that the human protein referred to
     herein as INSP076 protein is a member of the cytokine receptor-type I
     family (hematopoietin receptor superfamily). Preferably, INSP076
     functions as an IL-9 receptor or an IL-9 receptor-like protein. The
     INSP076 protein does not possess a transmembrane domain and accordingly
     the INSP076 protein is a potential soluble receptor. It is believed that the
     INSP076 protein may function as an IL-9 antagonist.
```

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,

AB

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RE.CNT 5

ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> save temp 128 brdstsubgen/a
ANSWER SET L28 HAS BEEN SAVED AS 'BRDSTSUBGEN/A'

=> save temp all niddmsrch/l L# LIST L1-L28 HAS BEEN SAVED AS 'NIDDMSRCH/L'

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COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION 21.69 588.62 FULL ESTIMATED COST DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION -2.92 -10.95CA SUBSCRIBER PRICE

SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 12:32:08 ON 26 APR 2005

Connecting via Winsock to STN

Welcome to STN International! Enter x:x

LOGINID: SSSPTA1623PAZ

PASSWORD:

* * * * * RECONNECTED TO STN INTERNATIONAL * * * * * * * SESSION RESUMED IN FILE 'CAPLUS' AT 13:08:50 ON 26 APR 2005 FILE 'CAPLUS' ENTERED AT 13:08:50 ON 26 APR 2005 COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

COST IN U.S COLLARS FULL ESTIMATED COST	SINCE FILE ENTRY 21.69	TOTAL SESSION 588.62
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) CA SUBSCRIBER PRICE	SINCE FILE ENTRY -2.92	TOTAL SESSION -10.95
=> file reg COST IN U.S. DOLLARS FULL ESTIMATED COST	SINCE FILE ENTRY 21.69	TOTAL SESSION 588.62
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) CA SUBSCRIBER PRICE	SINCE FILE ENTRY -2.92	TOTAL SESSION -10.95

FILE 'REGISTRY' ENTERED AT 13:08:57 ON 26 APR 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0 DICTIONARY FILE UPDATES: 25 APR 2005 HIGHEST RN 849177-50-0

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 18, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

****************** * The CA roles and document type information have been removed from * * the IDE default display format and the ED field has been added, * effective March 20, 2005. A new display format, IDERL, is now * available and contains the CA role and document type information. * *****

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

```
=> e 1,4-Benzenediacetic acid, diethyl ester/cn
                   1,4-BENZENEDIACETIC ACID, DIETHENYL ESTER, POLYMER WITH 1,4-
                   BUTANEDIOL/CN
                   1,4-BENZENEDIACETIC ACID, DIETHENYL ESTER, POLYMER WITH 1,6-
E2
             1
                   HEXANEDIOL/CN
E3
             1 --> 1,4-BENZENEDIACETIC ACID, DIETHYL ESTER/CN
                   1,4-BENZENEDIACETIC ACID, DIETHYL ESTER, COMPD. WITH 1-(ACET
E4
                   YLOXY)-N,N,N-TRIMETHYLMETHANAMINIUM SALT WITH 2,4,6-TRINITRO
                   PHENOL (1:1:1)/CN
                   1,4-BENZENEDIACETIC ACID, DIETHYL ESTER, COMPD. WITH N,N,N-T
E5
             1
                   RIMETHYLMETHANAMINIUM SALT WITH 2,4,6-TRINITROPHENOL (1:1:1)
                   /CN
                   1,4-BENZENEDIACETIC ACID, DIHEPTYL ESTER/CN
E6
             1
                   1,4-BENZENEDIACETIC ACID, DIHEXYL ESTER/CN
E7
             1
             1
                   1,4-BENZENEDIACETIC ACID, DIHYDRAZIDE, POLYMER WITH 1,4-BENZ
F.8
                   ENEDICARBONYL DICHLORIDE/CN
                   1,4-BENZENEDIACETIC ACID, DIMETHYL ESTER/CN
             1
E9
E10
             1
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                   -TRIMETHYL-1,6-HEXANEDIAMINE/CN
E11
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                   1,4-BENZENEDIACETIC ACID, DIMETHYL ESTER, POLYMER WITH DIMET
                   HYL 1,4-BENZENEDICARBOXYLATE AND C,C,C-TRIMETHYL-1,6-HEXANED
                   IAMINE/CN
                   1,4-BENZENEDIACETIC ACID, DINONYL ESTER/CN
E12
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=> e3
             1 "1,4-BENZENEDIACETIC ACID, DIETHYL ESTER"/CN
L29
=> d 129
    ANSWER 1 OF 1 REGISTRY COPYRIGHT 2005 ACS on STN
     36076-26-3 REGISTRY
```

RN

ED Entered STN: 16 Nov 1984

1,4-Benzenediacetic acid, diethyl ester (9CI) (CA INDEX NAME) OTHER CA INDEX NAMES:

p-Benżenediacetic acid, diethyl ester (6CI) OTHER NAMES:

CN NSC 139681

CN p-Phenylenediacetic acid diethyl ester

FS 3D CONCORD MF C14 H18 O4

CI COM

LC STN Files: BEILSTEIN*, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, TOXCENTER, USPAT2, USPATFULL (*File contains numerically searchable property data)

$$\begin{array}{c} \circ \\ \parallel \\ \text{CH}_2\text{-}\text{C}\text{-}\text{OEt} \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

20 REFERENCES IN FILE CA (1907 TO DATE)
20 REFERENCES IN FILE CAPLUS (1907 TO DATE).
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

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SESSION WILL BE HELD FOR 60 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 13:09:43 ON 26 APR 2005